```
Connecting via Winsock to STN
Welcome to STN International! Enter x:x
LOGINID:ssspat1756mja
PASSWORD:
LOGINID/PASSWORD REJECTED
The loginid and/or password sent to STN were invalid.
You either typed them incorrectly, or line noise may
have corrupted them.
Do you wish to retry the logon?
Enter choice (y/N):
Do you wish to use the same loginid and password?
Enter choice (y/N):
Enter new loginid (or press [Enter] for ssspat1756mja):
Enter new password:
LOGINID:
LOGINID: ssspta1756mja
PASSWORD:
TERMINAL (ENTER 1, 2, 3, OR ?):2
                      Welcome to STN International
 NEWS
                  Web Page URLs for STN Seminar Schedule - N. America
 NEWS 2
                  "Ask CAS" for self-help around the clock
 NEWS
      3 JUL 20 Powerful new interactive analysis and visualization software,
                  STN AnaVist, now available
      4 AUG 11 STN AnaVist workshops to be held in North America
 NEWS
      5 AUG 30 CA/Caplus -Increased access to 19th century research documents
 NEWS
      6 AUG 30 CASREACT - Enhanced with displayable reaction conditions
 NEWS
      7 SEP 09 ACD predicted properties enhanced in REGISTRY/ZREGISTRY
 NEWS
 NEWS
      8 OCT 03 MATHDI removed from STN
 NEWS
      9 OCT 04 CA/CAplus-Canadian Intellectual Property Office (CIPO) added
                  to core patent offices
                  STN AnaVist workshops to be held in North America
 NEWS 10
        OCT 06
         OCT 13
                 New CAS Information Use Policies Effective October 17, 2005
 NEWS 11
 NEWS 12 OCT 17
                 STN(R) AnaVist(TM), Version 1.01, allows the export/download
                  of CAplus documents for use in third-party analysis and
                  visualization tools
 NEWS 13
         OCT 27
                 Free KWIC format extended in full-text databases
 NEWS 14
         OCT 27 DIOGENES content streamlined
 NEWS 15 OCT 27 EPFULL enhanced with additional content
              JUNE 13 CURRENT WINDOWS VERSION IS V8.0, CURRENT
 NEWS EXPRESS
              MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
               AND CURRENT DISCOVER FILE IS DATED 13 JUNE 2005
 NEWS HOURS
               STN Operating Hours Plus Help Desk Availability
 NEWS INTER
               General Internet Information
 NEWS LOGIN
               Welcome Banner and News Items
 NEWS PHONE
              Direct Dial and Telecommunication Network Access to STN
 NEWS WWW
               CAS World Wide Web Site (general information)
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* STN
                                 Columbus
FILE 'HOME' ENTERED AT 08:49:00 ON 14 NOV 2005
=> file caplus
                                                  SINCE FILE
COST IN U.S. DOLLARS
                                                                  TOTAL
                                                       ENTRY
                                                                SESSION
                                                        0.21
                                                                   0.21
FULL ESTIMATED COST
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                                VOL 143 ISS 21
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                                (20051113/ED)
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=> s (nlo or nonlinear or (non(3a)linear) or hyperpolariz?)
          5079 NLO
             8 NLOS
          5082 NLO
                 (NLO OR NLOS)
        163729 NONLINEAR
             2 NONLINEARS
        163731 NONLINEAR
                 (NONLINEAR OR NONLINEARS)
        755839 NON
            34 NONS
        755866 NON
                 (NON OR NONS)
        556631 LINEAR
            55 LINEARS
        556655 LINEAR
                 (LINEAR OR LINEARS)
         20423 NON (3A) LINEAR
         19775 HYPERPOLARIZ?
        197907 (NLO OR NONLINEAR OR (NON(3A)LINEAR) OR HYPERPOLARIZ?)
=> s (((charge1 or hole1)(3a)(transport? or generat?)) or carbazole or vinylcarbazole or polyvinyl
1) (3A) (TRANSPORT? IS NOT A RECOGNIZED COMMAND
The previous command name entered was not recognized by the system.
For a list of commands available to you in the current file, enter
"HELP COMMANDS" at an arrow prompt (=>).
=> s (((charge or hole)(3a)(transport? or generat?)) or carbazole or vinylcarbazole or polyvinylca
        484684 CHARGE
         62432 CHARGES
        519427 CHARGE
                 (CHARGE OR CHARGES)
```

188697 HOLE 110661 HOLES 253252 HOLE

752033 TRANSPORT? 925147 GENERAT?

(HOLE OR HOLES)

```
38473 (CHARGE OR HOLE) (3A) (TRANSPORT? OR GENERAT?)
         15956 CARBAZOLE
          2128 CARBAZOLES
         16513 CARBAZOLE
                 (CARBAZOLE OR CARBAZOLES)
          5767 VINYLCARBAZOLE
           122 VINYLCARBAZOLES
          5783 VINYLCARBAZOLE
                 (VINYLCARBAZOLE OR VINYLCARBAZOLES)
          2223 POLYVINYLCARBAZOLE
            34 POLYVINYLCARBAZOLES
          2250 POLYVINYLCARBAZOLE
                 (POLYVINYLCARBAZOLE OR POLYVINYLCARBAZOLES)
         57950 (((CHARGE OR HOLE)(3A)(TRANSPORT? OR GENERAT?)) OR CARBAZOLE OR
L2
               VINYLCARBAZOLE OR POLYVINYLCARBAZOLE)
=> s 11 and 12
L3
          1293 L1 AND L2
=> s (photochrom? or diarylethane or diheteroarylethane or spiropyran or fulgide or leuco or photo
         12436 PHOTOCHROM?
           178 DIARYLETHANE
           234 DIARYLETHANES
           340 DIARYLETHANE
                 '(DIARYLETHANE OR DIARYLETHANES)
             0 DIHETEROARYLETHANE
          1635 SPIROPYRAN
           993 SPIROPYRANS
          1990 SPIROPYRAN
                 (SPIROPYRAN OR SPIROPYRANS)
           480 FULGIDE
           317 FULGIDES
           568 FULGIDE
                 (FULGIDE OR FULGIDES)
         11290 LEUCO
             7 LEUCOS
         11295 LEUCO
                 (LEUCO OR LEUCOS)
         10084 PHOTOISOMER?
        345680 ISOMER?
L4
        369617 (PHOTOCHROM? OR DIARYLETHANE OR DIHETEROARYLETHANE OR SPIROPYRAN
                OR FULGIDE OR LEUCO OR PHOTOISOMER? OR ISOMER?)
=> s 13 and 14
            27 L3 AND L4
=> d all 1-3
     ANSWER 1 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
L5
AN
     2005:1086283 CAPLUS
DN
     143:325922
     Entered STN:
ED
                   11 Oct 2005
     Dependence of the Two-Photon Absorption Cross Section on the Conjugation
     of the Phenylacetylene Linker in Dipolar Donor-Bridge-Acceptor
     Chromophores
ΑU
     Lee, Soohyun; Thomas, K. R. Justin; Thayumanavan, S.; Bardeen, Christopher
CS
     Departments of Chemistry, University of California, Riverside, CA, 92521,
SO
     Journal of Physical Chemistry A (2005), 109(43), 9767-9774
     CODEN: JPCAFH; ISSN: 1089-5639
     American Chemical Society
PB
DT
     Journal
LA
     English
CC
     22-9 (Physical Organic Chemistry)
     Section cross-reference(s): 73
AB
           ***nonlinear***
                            optical properties of four
                                                           ***isomeric***
     dipolar two-photon chromophores are compared. The chromophores consist of
         ***carbazole***
                          electron donor coupled to a naphthalimide electron
     acceptor by a phenylacetylene bridge. By variation of the connectivity of
     the bridge at the Ph groups, four compds. with 0, 1, and 2 meta linkages
     are synthesized. The linear and
                                       ***nonlinear***
                                                           optical properties of
```

these compds. are measured. Despite similar linear absorption cross sections, the two-photon absorption cross section .delta. of the all-meta compd. is almost a factor of 10 lower than the all-para compd. By taking the detailed mol. conformations into account in order to calc. accurate dipole moment changes, we find that the decrease in .delta. results largely from the decreased charge transfer ability with increasing no. of meta linkages. We find that a two-state model can be used to predict semiquant. the obsd. trend in .delta. on the basis of the linear optical properties of the mols. This work illustrates the dramatic effect the ground-state polarizability can have on the \*\*\*nonlinear\*\*\* response of org. compds. and also provides a way to quantify the ability of meta linkages to inhibit charge transfer in their ground-state configuration. two photon absorption conjugation phenylacetylene linker dipolar chromophore

- ST
- UV absorption IT

(UV-visible; dependence of two-photon absorption cross section on conjugation of phenylacetylene linker in dipolar donor-bridge-acceptor chromophores)

- Aromatic hydrocarbons, properties TT
  - RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (aryl alkynes, dipolar chromophores with phenylacetylene linker; dependence of two-photon absorption cross section on conjugation of phenylacetylene linker in dipolar donor-bridge-acceptor chromophores)
- IT
  - RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (aryl, dipolar chromophores with phenylacetylene linker; dependence of two-photon absorption cross section on conjugation of phenylacetylene linker in dipolar donor-bridge-acceptor chromophores)
- Fluorescence quenching IT

(by intramol. electron transfer in polar solvents; dependence of two-photon absorption cross section on conjugation of phenylacetylene linker in dipolar donor-bridge-acceptor chromophores)

Chromophores

Fluorescence

Molar absorptivity

\*\*\*Nonlinear\*\*\* optical materials

Solvent polarity effect

(dependence of two-photon absorption cross section on conjugation of phenylacetylene linker in dipolar donor-bridge-acceptor chromophores)

Dipole moment

(difference between ground and excited states; dependence of two-photon absorption cross section on conjugation of phenylacetylene linker in dipolar donor-bridge-acceptor chromophores)

IT Electron transfer

> (intramol.; dependence of two-photon absorption cross section on conjugation of phenylacetylene linker in dipolar donor-bridge-acceptor chromophores)

Two-photon absorption

; dependence of two-photon absorption cross section \*\*\*nonlinear\*\*\* on conjugation of phenylacetylene linker in dipolar donor-bridge-acceptor chromophores)

Molecular structure-property relationship

(optical; dependence of two-photon absorption cross section on conjugation of phenylacetylene linker in dipolar donor-bridge-acceptor chromophores)

IΤ Molecules

> (size; dependence of two-photon absorption cross section on conjugation of phenylacetylene linker in dipolar donor-bridge-acceptor chromophores)

\*\*\*Nonlinear\*\*\* optical absorption

(two-photon; dependence of two-photon absorption cross section on conjugation of phenylacetylene linker in dipolar donor-bridge-acceptor chromophores)

- Conjugation (bond)
  - (.pi.-; dependence of two-photon absorption cross section on conjugation of phenylacetylene linker in dipolar donor-bridge-acceptor chromophores)
- IT 865443-61-4, 3-Bromo-9-propyl-9H- \*\*\*carbazole\*\*\* 865443-57-8 RL: RCT (Reactant); RACT (Reactant or reagent)

(coupling with (trimethylsily1)acetylene; dependence of two-photon absorption cross section on conjugation of phenylacetylene linker in

```
dipolar donor-bridge-acceptor chromophores)
IT.
     865443-53-4P
                    865443-54-5P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (coupling with (trimethylsilyl)acetylene; dependence of two-photon
        absorption cross section on conjugation of phenylacetylene linker in
        dipolar donor-bridge-acceptor chromophores)
                                       591-18-4, 1-Bromo-3-iodobenzene
IT
     589-87-7, 1-Bromo-4-iodobenzene
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (coupling with alkyne; dependence of two-photon absorption cross
        section on conjugation of phenylacetylene linker in dipolar
        donor-bridge-acceptor chromophores)
IT
     865443-59-0P
                    865443-60-3P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (coupling with alkyne; dependence of two-photon absorption cross
        section on conjugation of phenylacetylene linker in dipolar
        donor-bridge-acceptor chromophores)
IT
     1066-54-2, (Trimethylsilyl)acetylene
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (coupling with arylbromide; dependence of two-photon absorption cross
        section on conjugation of phenylacetylene linker in dipolar
        donor-bridge-acceptor chromophores)
IT
     865443-55-6P
                    865443-56-7P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (coupling with arylbromide; dependence of two-photon absorption cross
        section on conjugation of phenylacetylene linker in dipolar
        donor-bridge-acceptor chromophores)
                    865443-58-9P
     865443-52-3P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (coupling with aryliodide; dependence of two-photon absorption cross
        section on conjugation of phenylacetylene linker in dipolar
        donor-bridge-acceptor chromophores)
IT
     865443-48-7P
                    865443-49-8P
                                   865443-50-1P
                                                  865443-51-2P
     RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
        (dependence of two-photon absorption cross section on conjugation of
        phenylacetylene linker in dipolar donor-bridge-acceptor chromophores)
RE.CNT
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    ANSWER 2 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
ΑN
    2005:951479 CAPLUS
ED
    Entered STN:
                  31 Aug 2005
    Light-induced phenomena in polymeric thin films
TI
AU
    Nespurek, S.; Pospisil, J.
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- Institute of Macromolecular Chemistry, Academy of Sciences of the Czech Republic, Prague, 162 06, Czech Rep.
- Journal of Optoelectronics and Advanced Materials (2005), 7(3), 1157-1168 CODEN: JOAMF2; ISSN: 1454-4164
- PB National Institute for Optoelectronics
- DT Journal
- LA English
- ĊC 74 (Radiation Chemistry, Photochemistry, and Photographic and Other Reprographic Processes)
- Photoprocesses in polymers are generally divided into two groups: (i) Photochem. processes resulting in a permanent chem. alternation of a macromol., like bond scission, crosslinking and oxidn., (ii) Phys. (reversible) processes involving the distribution, relocation and ultimate fate of the excess energy assocd. with an excited mol., following absorption of a photon, like photoluminescence, phosphorescence, photoinduced electron transfer and formation of charge-transfer states (excitons). The excitation is very often accompanied by charge redistribution and change of mol. conformation. An extreme case is the \*\*\*photochromic\*\*\* effect - photoreversible reaction of two forms of a single mol. The changes in the mol. conformations lead very often to the formation of metastable electronic states. Similarly to semiconductors, electron-hole pairs or ion-pairs can be generated by the action of light in conjugated polymers. Their dissocn. leads to the \*\*\*generation\*\*\* of free \*\*\*charge\*\*\* carriers. Photoconductive polymers with optical \*\*\*non\*\*\* - \*\*\*linear\*\*\* activity can be used for photorefractive and holog. memories.

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     ANSWER 3 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
     2004:296729 CAPLUS
ΑN
DN
     142:45454
     Entered STN: 12 Apr 2004
     Photorefractive gratings in DRY-doped hybrid sol-gel films
     Raschella, R.; Marino, I.-G.; Lottici, P. P.; Bersani, D.; Lorenzi, A.;
     Istituto Nazionale per la Fisica della Materia (INFM) and Dipartimento di
     Fisica, Raman-Exafs Laboratory, Universita di Parma, Parma, 43100, Italy
     Optical Materials (Amsterdam, Netherlands) (2004), 25(4), 419-423
     CODEN: OMATET; ISSN: 0925-3467
PB
    Elsevier Science B.V.
DT
    Journal
LA
     English
     73-11 (Optical, Electron, and Mass Spectroscopy and Other Related
CC
     Properties)
    Photorefractive gratings were obtained with 632.8 nm writing beams in
AB
     org.-inorg. SiO2-based films. The hybrid glass was prepd. by a sol-gel
     technique, starting from org. Si precursors, and contains Disperse Red 1
     (DR1), ***carbazole***
                               units and 2,4,7-trinitro-9-fluorenone (TNF).
     The photorefractive gain, which was found unexpectedly even without poling
     field, was detd. through an asym. energy exchange by two-beam coupling
     measurements. The effects of the polarization of the writing beams and of
                             ***photoisomerizing*** radiation during the
     a circularly polarized
     erasure of the grating were interpreted in terms of an orientational
     contribution to the grating formation.
ST
     photorefractive grating DR1 doped hybrid sol gel film
IT
       ***Nonlinear***
                        optical properties
        (beam coupling; photorefractive gratings in DRY-doped hybrid sol-gel
        films)
IT
     Photorefractive effect
        (gain; photorefractive gratings in DRY-doped hybrid sol-gel films)
IT
     Energy transfer
     Optical gain
     Photorefractive gratings
     Sol-gel processing
        (photorefractive gratings in DRY-doped hybrid sol-gel films)
IT
     2872-52-8, Disperse Red 1
     RL: OCU (Occurrence, unclassified); PRP (Properties); OCCU (Occurrence)
        (photorefractive gratings in DRY-doped hybrid sol-gel films)
                     919-30-2, 3-Aminopropyltriethoxysilane 24801-88-5,
     78-10-4, TEOS
                                           73500-82-0,
                                                         ***Carbazole***
     3-(Triethoxysilyl)propyl-isocyanate
     -9carbonyl chloride
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (photorefractive gratings in DRY-doped hybrid sol-gel films)
RE.CNT
       18
             THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE
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=> d al 4-27
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ALL ----- BIB, AB, IND, RE
APPS ----- AI, PRAI
BIB ------ AN, plus Bibliographic Data and PI table (default)
CAN ----- List of CA abstract numbers without answer numbers
CBIB ----- AN, plus Compressed Bibliographic Data
DALL ----- ALL, delimited (end of each field identified)
DMAX ----- MAX, delimited for post-processing
FAM ----- AN, PI and PRAI in table, plus Patent Family data
FBIB ----- AN, BIB, plus Patent FAM
IND ----- Indexing data
IPC ----- International Patent Classifications
MAX ----- ALL, plus Patent FAM, RE
PATS ----- PI, SO
SAM ----- CC, SX, TI, ST, IT
SCAN ----- CC, SX, TI, ST, IT (random display, no answer numbers;
             SCAN must be entered on the same line as the DISPLAY,
             e.g., D SCAN or DISPLAY SCAN)
STD ----- BIB, IPC, and NCL
IABS ----- ABS, indented with text labels
IALL ----- ALL, indented with text labels
IBIB ----- BIB, indented with text labels
IMAX ----- MAX, indented with text labels
ISTD ----- STD, indented with text labels
OBIB ----- AN, plus Bibliographic Data (original)
OIBIB ----- OBIB, indented with text labels
SBIB ----- BIB, no citations
SIBIB ----- IBIB, no citations
HIT ----- Fields containing hit terms
HITIND ----- IC, ICA, ICI, NCL, CC and index field (ST and IT)
             containing hit terms
HITRN ----- HIT RN and its text modification
HITSTR ----- HIT RN, its text modification, its CA index name, and
             its structure diagram
HITSEQ ----- HIT RN, its text modification, its CA index name, its
             structure diagram, plus NTE and SEQ fields
FHITSTR ---- First HIT RN, its text modification, its CA index name, and
             its structure diagram
FHITSEQ ---- First HIT RN, its text modification, its CA index name, its
             structure diagram, plus NTE and SEQ fields
KWIC ----- Hit term plus 20 words on either side
OCC ----- Number of occurrence of hit term and field in which it occurs
To display a particular field or fields, enter the display field
codes. For a list of the display field codes, enter HELP DFIELDS at
an arrow prompt (=>). Examples of formats include: TI; TI,AU; BIB,ST;
```

TI, IND; TI, SO. You may specify the format fields in any order and the

information will be displayed in the same order as the format

specification.

All of the formats (except for SAM, SCAN, HIT, HITIND, HITRN, HITSTR, FHITSTR, HITSEQ, FHITSEQ, KWIC, and OCC) may be used with DISPLAY ACC to view a specified Accession Number. ENTER DISPLAY FORMAT (BIB):d all 4-47 'D' IS NOT A VALID FORMAT FOR FILE 'CAPLUS' The following are valid formats: ABS ----- GI and AB ALL ----- BIB, AB, IND, RE APPS ----- AI, PRAI BIB ----- AN, plus Bibliographic Data and PI table (default) CAN ----- List of CA abstract numbers without answer numbers CBIB ----- AN, plus Compressed Bibliographic Data DALL ----- ALL, delimited (end of each field identified) DMAX ----- MAX, delimited for post-processing FAM ----- AN, PI and PRAI in table, plus Patent Family data FBIB ----- AN, BIB, plus Patent FAM IND ----- Indexing data IPC ----- International Patent Classifications MAX ----- ALL, plus Patent FAM, RE PATS ----- PI, SO SAM ----- CC, SX, TI, ST, IT SCAN ----- CC, SX, TI, ST, IT (random display, no answer numbers; SCAN must be entered on the same line as the DISPLAY, e.g., D SCAN or DISPLAY SCAN) STD ----- BIB, IPC, and NCL IABS ----- ABS, indented with text labels IALL ----- ALL, indented with text labels IBIB ----- BIB, indented with text labels IMAX ----- MAX, indented with text labels ISTD ----- STD, indented with text labels OBIB ----- AN, plus Bibliographic Data (original) OIBIB ----- OBIB, indented with text labels SBIB ----- BIB, no citations SIBIB ----- IBIB, no citations HIT ----- Fields containing hit terms HITIND ----- IC, ICA, ICI, NCL, CC and index field (ST and IT) containing hit terms HITRN ----- HIT RN and its text modification HITSTR ----- HIT RN, its text modification, its CA index name, and its structure diagram HITSEQ ----- HIT RN, its text modification, its CA index name, its structure diagram, plus NTE and SEQ fields FHITSTR ---- First HIT RN, its text modification, its CA index name, and its structure diagram FHITSEQ ---- First HIT RN, its text modification, its CA index name, its structure diagram, plus NTE and SEQ fields KWIC ----- Hit term plus 20 words on either side OCC ----- Number of occurrence of hit term and field in which it occurs To display a particular field or fields, enter the display field codes. For a list of the display field codes, enter HELP DFIELDS at an arrow prompt (=>). Examples of formats include: TI; TI, AU; BIB, ST; TI, IND; TI, SO. You may specify the format fields in any order and the information will be displayed in the same order as the format specification. All of the formats (except for SAM, SCAN, HIT, HITIND, HITRN, HITSTR, FHITSTR, HITSEQ, FHITSEQ, KWIC, and OCC) may be used with DISPLAY ACC

ANSWER 4 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

Org.-inorg. films based on SiO2, contg. Disperse Red 1 (DR1),

\*\*\*carbazole\*\*\* units and 2,4,7-trinitro-9-fluorenone (TNF), have been prepd. by a sol-gel technique. Diffraction gratings have been produced through different effects: photoinduced. . . coupling (2BC)

to view a specified Accession Number.

ENTER DISPLAY FORMAT (BIB): kwic

measurements. The effect of the polarization of the writing beams on the grating and of a circularly polarized \*\*\*photoisomerizing\*\*\* during grating erasure has been interpreted in terms of an orientation contribution to the grating formation. \*\*\*Isomerization\*\*\* (cis-trans, photochem.; recording of holog. gratings in hybrid org.-inorg. films based on SiO2 sol-gel films contg. Disperse Red 1 and \*\*\*carbazole\*\*\* units and trinitrofluorenone) (photoinduced; recording of holog. gratings in hybrid org.-inorg. films based on SiO2 sol-gel films contg. Disperse Red 1 and \*\*\*carbazole\*\*\* units and trinitrofluorenone) Holographic diffraction gratings Holographic recording materials Hybrid organic-inorganic materials Molecular orientation Photorefractive effect Sol-gel processing (recording of holog. gratings in hybrid org.-inorg. films based on SiO2 sol-gel films contg. Disperse Red 1 and \*\*\*carbazole\*\*\* units and trinitrofluorenone) \*\*\*Nonlinear\*\*\* optical properties (two-beam-coupling; recording of holog. gratings in hybrid org.-inorg. films based on SiO2 sol-gel films contg. Disperse Red 1 and \*\*\*carbazole\*\*\* units and trinitrofluorenone) 7647-01-0, Hydrochloric acid, uses RL: CAT (Catalyst use); USES (Uses) (recording of holog. gratings in hybrid org.-inorg. films based on SiO2 sol-gel films contg. Disperse Red 1 and \*\*\*carbazole\*\*\* units and trinitrofluorenone) 685091-04-7 RL: PEP (Physical, engineering or chemical process); PYP (Physical process); PROC (Process) (recording of holog, gratings in hybrid org.-inorg, films based on SiO2 sol-gel films contg. Disperse Red 1 and \*\*\*carbazole\*\*\* trinitrofluorenone) 919-30-2, 3-Aminopropyltriethoxysilane 2872-52-8, Disperse Red 1 24801-88-5, 3-(Triethoxysilyl)propylisocyanate 73500-82-0, \*\*\*Carbazole\*\*\* -9-carbonyl chloride RL: PEP (Physical, engineering or chemical process); PYP (Physical process); RCT (Reactant); PROC (Process); RACT (Reactant or reagent) (recording of holog. gratings in hybrid org.-inorg. films based on SiO2 sol-gel films contg. Disperse Red 1 and \*\*\*carbazole\*\*\* trinitrofluorenone) 129-79-3, 2,4,7-Trinitro-9-fluorenone RL: PEP (Physical, engineering or chemical process); PYP (Physical process); PROC (Process) (sensitizer; recording of holog. gratings in hybrid org.-inorg. films based on SiO2 sol-gel films contg. Disperse Red 1 and \*\*\*carbazole\*\*\* units and trinitrofluorenone) ANSWER 5 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN . realized in the case where a doping of polymers by low-mol.-wt. dyed impurities does not essentially distort spectral, luminescent and optical properties of the individual components of \*\*\*nonlinear\*\*\* polymer compn. The influence of polymer nature and chem. constitution of . . was considered. The main paths of energy degrdn. of electronic excitation in such materials - internal conversion, electron \*\*\*photoisomerization\*\*\* transfer and - were analyzed. The role of dyes in the process of generation, in recombination and in the carriage of charges. 25067-59-8, \*\*\*Polyvinylcarbazole\*\*\* 55774-96-4, Poly-N-epoxypropylcarbazole RL: DEV (Device component use); PRP (Properties); USES (Uses) (mol. engineering of dye-doped polymers for optoelectronics) ANSWER 6 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN CdSe/CdS core/shell, tri-n-octylphosphine oxide passivated, quantum dots are used to sensitize a photorefractive polymer composite. The composite also consists of poly(N- \*\*\*vinylcarbazole\*\*\* ) as the nominally

\*\*\*transporting\*\*\* matrix and an electrooptic

chromophore. The efficacy of sensitization and consequent photorefractive

IT

IT

IT

TT

IT

L5

ΑB

L5

AB

\*\*\*charge\*\*\*

```
performance is investigated using transmission spectroscopy and
     ellipsometry,.
IT
       ***Isomerization***
       (cis-trans, photochem.; photorefractive poly( ***vinylcarbazole***
        )-based composite contg. (ethylhexyloxydimethylnitrophenylazo)benzene
        sensitized by octylphosphine oxide passivated CdSe/CdS quantum dots in
        relation to)
IT
     Degenerate four wave mixing
     Ellipsometry
     Holographic diffraction gratings
     Nanoparticles
     Photorefractive effect
     Photorefractive materials
     Quantum dot devices
     Refractive index
     Space charge
        (photorefractive poly( ***vinylcarbazole*** )-based composite contg.
        (ethylhexyloxydimethylnitrophenylazo)benzene sensitized by
        octylphosphine oxide passivated CdSe/CdS quantum dots)
     Holographic recording materials
IT
        (photorefractive poly( ***vinylcarbazole*** )-based composite contg.
        (ethylhexyloxydimethylnitrophenylazo) benzene sensitized by
        octylphosphine oxide passivated CdSe/CdS quantum dots in relation to)
IT
     Birefringence
        (transient; photorefractive poly( ***vinylcarbazole*** )-based
        composite contg. (ethylhexyloxydimethylnitrophenylazo)benzene
        sensitized by octylphosphine oxide passivated CdSe/CdS quantum dots)
IT
       ***Nonlinear***
                       optical properties
        (two-beam-coupling; photorefractive poly( ***vinylcarbazole***
        )-based composite contg. (ethylhexyloxydimethylnitrophenylazo)benzene
        sensitized by octylphosphine oxide passivated CdSe/CdS quantum dots)
IT
     78-50-2, Tri-n-octylphosphine oxide 1306-23-6, Cadmium sulfide,
                  1306-24-7, Cadmium selenide, properties
     properties
                                                             25067-59-8, Poly(N-
       ***vinylcarbazole*** ) 176681-76-8, EHDNPB
     RL: PEP (Physical, engineering or chemical process); PRP (Properties); PYP
     (Physical process); PROC (Process)
        (photorefractive poly( ***vinylcarbazole*** )-based composite contg.
        (ethylhexyloxydimethylnitrophenylazo)benzene sensitized by
        octylphosphine oxide passivated CdSe/CdS quantum dots)
     ANSWER 7 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
L5
     The mechanism of optical nonlinearity of new polymer composites based on a
AΒ
     conducting polymer [poly(9- ***vinylcarbazole*** )] and fullerenes C70 and C60 was studied both exptl. and theor. The ***nonlinear***
     -optical studies of self-action and coupling of two 633-nm beams from a
     He-Ne laser were performed, and variations in the absorption. . . in
     polarizabilities of fullerene mols. and their anion radicals, which are
     formed upon absorption of photons and charge transfer by poly(9-
       ***vinylcarbazole*** ) mols.
ST
     fullerene polyvinyl
                          ***carbazole***
                                                ***nonlinear***
                                                                  optical
     property nanocomposite
IT
     Conducting polymers
     Nanocomposites
         ***Nonlinear***
                          optical susceptibility
         ***Photochromism***
     Photorefractive effect
        (optical nonlinearity of fullerene-doped polymer nanocomposites)
TT
     25067-59-8, Poly(9- ***vinylcarbazole*** ) 99685-96-8, Fullerene (C60)
     115383-22-7, Fullerene C70
     RL: PRP (Properties)
        (optical nonlinearity of fullerene-doped polymer nanocomposites)
L5
     ANSWER 8 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
ΤI
     Synthesis, properties and
                                ***photochromism***
                                                       of novel charge transfer
     compounds with Keggin anions and protonated 2,2'-biquinoline
AΒ
     Several novel compds. with protonated 2,2'-biquinoline (biqui) and Keggin
     polyoxoanions (.alpha.- ***isomers*** ), (Hbiqui)m[XM12O40].cntdot.n(sol
     V), X = P (m = 3), Si (m = 4); M = Mo, W; n = 0, 3; solv.
     disordered 2,2'-biquinoline mols. and .alpha.-[SiW12O40]4- anions.
     Photosensitivity to sunlight and to W-lamp visible light was assessed for
                    ***Photochromic***
                                        properties were found for solids with
```

[XMo12O40]m- Keggin anions. Redn. of these anions upon irradn. was obsd.

```
by diffuse reflectance. . . sunlight, the extent of anion redn.
followed the order (Hbiqui)3[PMo12040].cntdot.3DMF>(Hbiqui)4[SiMo12040].cn
tdot.3DMF>(Hbiqui)3[PMo12O40]>(Hbiqui)4[SiMo12O40].cntdot.4H2O. The Kurtz
powder test was used to evaluate the 2nd-order
                                                ***nonlinear***
                                                                  optical
properties of the prepd. compds. (Hbiqui)4[SiW12040].cntdot.3H20
originated a 2nd harmonic generation signal with intensity .apprx.15% that
of urea for.
Keggin biquinolinium charge transfer prepn UV;
                                                ***nonlinear***
                                                                  optical
property tungstosilicate biquinolinium charge transfer;
  ***photochromism***
                       molybdophosphate molybdosilicate biquinolinium
charge transfer; biquinolinium molybdophosphate molybdosilicate
tungstophosphate tungstosilicate charge transfer prepn UV
Heteropoly acids
RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
                                   ***photochromism***
                                                          of charge
   (Keggin; prepn., properties and
   transfer compds. with Keggin anions and 2,2'-biquinolinium)
  ***Photochromism***
   (in molybdosilicate and molybdophosphate biquinolinium charge transfer
   complexes)
                 ***generation***
Second-harmonic
   (of tungstosilicate biquinolinium ***charge***
                                                     transfer complex)
Charge transfer complexes
RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
   (prepn., properties and
                            ***photochromism*** of charge transfer
   compds. with Keggin anions and 2,2'-biquinolinium)
              503014-09-3P 503014-15-1P 503014-17-3P
503014-07-1P
RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
   (prepn., UV-visible spectra and ***photochromism*** )
ANSWER 9 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
The origin of the optical nonlinearity of an org. polymeric compn.
including fullerene C70 and poly(9- ***vinylcarbazole*** ) (PVK) was
investigated by two-beam coupling and self-action of a cw laser beam at
633 nm, and spectroscopically. The local.
  ***photochromism***
                       optical nonlinearity ***polyvinylcarbazole***
fullerene
  ***Nonlinear***
                   optical properties
    ***Photochromism***
Polarizability
   (optical nonlinearity of C70-sensitized photorefractive compns. caused
   by prodn. of anion-radicals)
25067-59-8, Poly(9- ***vinylcarbazole*** )
                                              115383-22-7, C70 Fullerene
RL: PRP (Properties)
   (optical nonlinearity of C70-sensitized photorefractive compns. caused
   by prodn. of anion-radicals)
ANSWER 10 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
Multi-grating in photorefractive composites containing
                                                        ***nonlinear***
chromophore azo-dye 1-n-butoxyl-2,5-dimethyl-4-(4'-nitrophenylazo)benzene
Distinct electroabsorption grating and
                                       ***photoisomerization***
grating as well as photorefractive grating were obsd. in the composite
consisting of 1-n-butoxyl-2,5- dimethyl-4-(4'-nitrophenylazo)benzene
(BDMNPAB): poly(N- ***vinylcarbazole*** ) (PVK): 2,4,7-trinitro-9-
fluorenone (TNF) in a wt. ratio of 44:55:1. Based on the quick
translation technique of two-beam coupling exptl. geometry, the authors
measured the electroabsorption grating to be 6-7 cm-1, and
  ***photoisomeric***
                       effect also brought an absorption grating of 2-3
         ***Photoisomeric***
                             effect caused index grating was sepd. from
the others by analyzing the dynamic behavior of the gratings, and an
amplitude. .
multi grating photorefractive composite
                                         ***nonlinear***
                                                          chromophore
azo dye; photorefractive grating
                                  ***photoisomerization***
electroabsorption two beam coupling
  ***Nonlinear***
                  optical properties
   (beam coupling; multi-grating in photorefractive composites contg.
     ***nonlinear***
                      chromophore azo-dye 1-n-butoxyl-2,5-dimethyl-4-(4'-
  nitrophenylazo)benzene)
Azo dyes
Electric field effects
Electrooptical absorption
Photorefractive gratings
   (multi-grating in photorefractive composites contg. ***nonlinear***
```

IT

IT

IT

IT

IT

L5

AB

```
chromophore azo-dye 1-n-butoxyl-2,5-dimethyl-4-(4'-
       nitrophenylazo)benzene)
IT
       ***Isomerization*** .
          ***photoisomerization*** ; multi-grating in photorefractive
                           ***nonlinear***
                                             chromophore azo-dye
       composites contg.
       1-n-butoxyl-2,5-dimethyl-4-(4'-nitrophenylazo)benzene)
     129-79-3, 2,4,7-Trinitro-9-fluorenone 25067-59-8, Poly(N-
IT
       ***vinylcarbazole*** )
                                210281-60-0
    RL: DEV (Device component use); PRP (Properties); USES (Uses)
                                                             ***nonlinear***
        (multi-grating in photorefractive composites contg.
       chromophore azo-dye 1-n-butoxyl-2,5-dimethyl-4-(4'-
       nitrophenylazo)benzene)
    ANSWER 11 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
L5
       ***Photoisomeric*** -chromophore-doped polymers for directional spatial
ΤI
    filtering by optical threshold processing
AB
    We demonstrate spatial filtering by using the azo-chromophore-based
       ***nonlinear***
                       optical effect of self-beam polarization modulation and
    optical threshold processing. The real-time image-processing scheme is
    described by use of an. . . system, Fourier-transform operation, and an
    azo-chromophore-doped film as a spatial filter. As Disperse Red 1 dye
     (DR1) doped in an N-poly( ***vinylcarbazole*** ) (PVK) film with
    N-ethylcarbazole (ECZ) of low glass-transition temp. exhibits
    self-polarization rotation after thin-film transmission, a linearly
    polarized probe beam.
                           . . of an original input image can be manipulated
    as they pass through the film. These effects are responsible for the
       ***photoisomerization*** of DR1 mols. in the low-glass-transition-temp.
    polymer matrix.
ST
       ***photoisomeric***
                            chromophore doped polymer; directional spatial
    filtering optical threshold processing
IT
       ***Isomerization***
                                 ***photoisomeric*** -chromophore-doped
        (cis-trans, photochem.;
       polymers for directional spatial filtering by optical threshold
       processing)
IT
    Memory devices
        ***Nonlinear*** optical materials
    Optical films
    Optical filters
    Optical modulation
    Spatial light modulators
          ***photoisomeric*** -chromophore-doped polymers for directional
       spatial filtering by optical threshold processing)
    86-28-2, N-Ethylcarbazole 2872-52-8, Disperse Red 1
IT
                                                            25067-59-8,
               ***carbazole*** )
    Poly(vinyl
    RL: PRP (Properties); TEM (Technical or engineered material use); USES
     (Uses)
          ***photoisomeric*** -chromophore-doped polymers for directional
       spatial filtering by optical threshold processing)
    ANSWER 12 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
1.5
ΤI
    Development of self adaptive image processing system by use of a
      ***photoisomeric*** organic dye film
    A review with 8 refs. We demonstrate spatial filtering using the
AB
                           ***nonlinear*** optical effect of the self-beam
    azo-chromophore-based
    polarization-modulation. The real-time image processing scheme is
    described using the optical configuration, composed of an.
    action-probe beam system. Fourier-transform operation and an
    azo-chromophore-doped film as a spatial filter. As Disperse Red 1 doped
    in the N- ***polyvinylcarbazole*** film with N-ethylcarbazole, of low
    glass-transition-temp., exhibits the self polarization-modulation after
    the thin film transmission, a linearly-polarized probe beam into.
    of an original input image can be manipulated as they pass through the
    film. This effect is responsible for the ***photoisomerization***
    DR1 mols. in the low-glass-transition-temp. polymer matrix.
    review self adaptive image processing; ***photoisomeric***
    film review
IT
    Imaging
    Optical films
    Optical filters
    Spatial light modulators
        (development of self adaptive image processing system by use of a
         ***photoisomeric*** org. dye film)
```

```
Memory effect
        (optical; development of self adaptive image processing system by use
              ***photoisomeric***
                                    org. dye film)
TΤ
      .***Isomerization***
          ***photoisomérization*** ; development of self adaptive image
       processing system by use of a ***photoisomeric***
                                                            org. dye film)
IT
    86-28-2, N-Ethylcarbazole
                              2872-52-8, C.I. Disperse Red 1
                                                                 25067-59-8,
    N- ***Polyvinylcarbazole***
    RL: PRP (Properties); TEM (Technical or engineered material use); USES
     (Uses)
        (development of self adaptive image processing system by use of a
         ***photoisomeric*** orq. dye film)
    ANSWER 13 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
L5
    sol gel org inorg optical properties; xerogel pyrromethene optical
st
    pumping; merocyanine spiro oxazine ***photochromic*** sol gel; laser
    written grating DR1 sol gel; photorefractive sol gel ***carbazole***
       ***Photochromism***
IT
        (and reverse- ***photochromism***
                                            of spiro-oxazine in sol-gel
       matrix)
      ***Nonlinear***
                       optical properties
        (two-beam coupling; in photorefractive sol-gel materials)
IT
    206867-72-3D, derivs.
                           206867-73-4D, derivs.
    RL: PEP (Physical, engineering or chemical process); PRP (Properties);
    PROC (Process)
          ***photochromism***
                                and reverse- ***photochromism***
       matrix)
                                     998-30-1, Triethoxysilane
IT
    78-08-0, Vinyl triethoxysilane
    Methyltriethoxysilane
    RL: PEP (Physical, engineering or chemical process); PROC (Process)
          ***photochromism*** and reverse- ***photochromism***
       spiro-oxazine in sol-gel matrix from)
IT
    86-74-8D, ***Carbazole*** , silicon alkoxide deriv.
                                                             2872-52-8D,
    Disperse red 1, silicon alkoxide deriv.
    RL: PEP (Physical, engineering or chemical process); PRP (Properties);
    PROC (Process)
        (photorefractive sol-gel materials using)
L5
    ANSWER 14 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
AΒ
       . . to their comparatively low glass-transition temps. Tg, an
    orientational enhancement of the photorefractive properties was obsd.
    Furthermore, the influence of
                                   ***photoisomerization***
                                                               (based on
     trans-cis-trans cycles) on the holog. properties could be detd. for the
    different chromophores. In addn., a class of fully functionalized
    polymers with azo chromophores and ***carbazole*** -units covalently
    attached to PMMA- and PU-backbones was synthesized. These systems show
     comparatively high glass transition temps. of more than 80 degree.C.
     third type of materials investigated is a glass of triphenylamine with
              ***carbazole*** and ***NLO*** -chromophore moieties. It
    has a glass transition temp. of 120.degree.C. For the high-Tg materials,
    poling procedures - essential for the photorefractive properties - could
    be monitored in-situ by second-harmonic generation. Abs. values for the
       ***nonlinear*** Pockels coeffs. .chi.(2)(-.omega.;.omega.,0) have been
    obtained by electro-optical measurements.
IT
       ***Isomerization***
          ***photoisomerization*** ; photorefractive polymers for real-time
       holog.)
L<sub>5</sub>
    ANSWER 15 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
ΤI
    Linear and second order ***nonlinear*** optical properties of novel
      ***photochromic*** materials
AB
    A new unique ***photochromic***
                                        material which is based on a
    reversible formation - cleavage of a C-C bond is described. The bicyclic
     (spiro) bindon deriv. (.lambda.max = 480 nm in toluene) undergoes
    photochem. and/or thermally induced ring opening to form the
                                                              ***isomer***
       ***isomer***
                    (.lambda.max= 640 nm in toluene). This
    form presents a conjugated donor-acceptor system and exhibits a
    considerable second-order optical nonlinearity as found by field induced
                                 ***Photochromic*** conversion was also
    SHG (EFISH) measurements.
    obsd. in the cryst. form indicated visually by a crystal red-to-green
    color change. The reversible ring opening - closure process in liq. and
```

IT

```
Optical and thermal switching and the
                                            ***NLO***
                                                      efficiency of the
     guest-host polymer system are described.
     bindon dimethylaminocinnamic aldehyde
                                            ***isomer***
       ***photochromic*** conversion;
                                         ***nonlinear***
                                                           optical property
     biindantrione dimethylaminocinnamic aldehyde; solvatochromism
       ***photochromism*** indan deriv polymer host; polymethylmethacrylate
                            ***NLO*** ; ***polyvinylcarbazole***
     host indantrione deriv
     indantrione deriv
                       ***NLO***
       ***Isomerization***
IT
        (cis-trans, photochem. and thermally induced; linear and second order
          ***nonlinear*** optical properties of biindantrione deriv.
          ***photochromic***
                             material in polymer hosts)
     Optical switching
IT
         ***Photochromic***
                             materials
         ***Photochromism***
     Second-order
                   ***nonlinear***
                                     optical properties
     Solvatochromism
        (linear and second order ***nonlinear***
                                                    optical properties of
       biindantrione deriv. ***photochromic*** material in polymer hosts)
IT
     56-23-5, Carbon tetrachloride, uses 67-66-3, Chloroform, uses
    DMSO, uses 71-43-2, Benzene, uses 75-05-8, Acetonitrile, uses
     75-09-2, uses 108-88-3, Toluene, uses
                                             110-82-7, Cyclohexane, uses
     123-91-1, Dioxane, uses 9011-14-7, PMMA 25067-59-8, Poly(
       ***vinylcarbazole*** )
     RL: NUU (Other use, unclassified); USES (Uses)
        (linear and second order ***nonlinear***
                                                    optical properties of
        biindantrione deriv. ***photochromic***
                                                   material in polymer hosts)
     197023-27-1P
                   197023-28-2P
IT
     RL: PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); PREP
     (Preparation); RACT (Reactant or reagent)
        (linear and second order ***nonlinear*** optical properties of
        biindantrione deriv.
                              ***photochromic***
                                                   material in polymer hosts)
IT
     1707-95-5, Bindon 6203-18-5, p-(Dimethylamino)cinnamic aldehyde
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (linear and second order ***nonlinear***
                                                    optical properties of
        biindantrione deriv.
                              ***photochromic***
                                                   material in polymer hosts)
     ANSWER 16 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
       ***Carbazole*** -based multifunctional molecules for photorefractive
ΤI
     applications
                        ***photochromic***
AB
     A multifunctional
                                            mols. consist of a
                       part with a ***nonlinear*** optical chromophore unit
       ***carbazole***
     attached via an alkyl link to the N atom of the ***carbazole***
     These compds. exhibit dual functions: photocond. and a second order
       ***nonlinear***
                        optical activity. The photocharge transport and the
     electrooptic functions are performed by two sep. constituents.
     Photorefractive effect of ethyl(hexylsulfonylphenylazophenyl)aminododecylc
     arbazole (I).
    photorefractive
                     ***carbazole***
                                        based multifunctional mol holog;
       ***nonlinear*** optical substituent ***carbazole*** deriv
    photorefractive
    Holography
     Photorefractive effect
     Photorefractive materials
        (photorefractive effect and dynamic holog. recording using
          ***carbazole*** -based multifunctional mols.)
       ***Nonlinear*** optical properties
        (two beam-coupling; of photorefractive
                                                ***carbazole*** -based
       multifunctional mols.)
     196201-41-9
     RL: PRP (Properties); TEM (Technical or engineered material use); USES
     (Uses)
        (photorefractive effect and dynamic holog, recording using
          ***carbazole*** -based multifunctional mols.)
IT
     129-79-3, 2,4,7-Trinitro-9-fluorenone
     RL: TEM (Technical or engineered material use); USES (Uses)
        (photorefractive effect and dynamic holog. recording using
          ***carbazole*** -based multifunctional mols.)
    ANSWER 17 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
L5
          . are unique to semiconductor nanoclusters. For example, this
```

polymeric solns. [PMMA and poly( \*\*\*vinylcarbazole\*\*\* )] was studied.

```
technique provided some useful kinetic and mechanistic information of size
quantization effects, ***nonlinear*** optical effects, electro and
                      effects, photocurrent
                                              ***qeneration***
  ***photochromic***
photocatalysis and ***charge*** rectification properties of
semiconductor nanoclusters. These properties are controlled by charge
sepn., charge trapping and heterogeneous charge transfer at the. .
Nanoparticles
    ***Nonlinear***
                     optical properties
Quantum size effect
Semiconductor materials
   (transient absorption spectroscopy of semiconductor nanoclusters)
ANSWER 18 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
Photorefractive polymers which incorporate azo-dyes as the
                                                             ***non***
  ***linear*** chromophore element, can be used not only for generating
gratings by the photorefractive effect, but also by
  ***photoisomerization***
                           of the azo-dye. In the latter mechanism,
repeated trans-cis ***isomerization*** causes the chromophore mols. to
become aligned at right angles to the laser polarization direction,
thereby making the material birefringent.. . diffraction efficiency
of the photorefractive gratings is a very sensitive function of the poling
field strength, while that of the ***photoisomerization*** gratings is
less so. In this work, we investigate the components diffracted from each
of these grating formed in a.
  ***Isomerization***
   ( ***photoisomerization*** ; multiple-grating formation in
   photorefractive polymers with azo-dye chromophores)
2872-52-8, Disperse REd1 25067-59-8, Poly(N- ***vinylcarbazole*** )
119516-33-5, 4-N, N-Diethylamino-(E)-cinnamonitrile
RL: PRP (Properties)
   (multiple-grating formation in photorefractive polymers with azo-dye
   chromophores)
ANSWER 19 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
  . . interfaces using second harmonic generation methods is described.
Among the topics discussed are the dynamics of photoinduced structure
changes, the ***transport***
                                     ***charge***
                                of
                                                     across an interface,
the rotational motions of interfacial mols., intermol. energy transfer
within the interface, interfacial photopolymn., and photoprocesses at.
Electron exchange and Charge transfer
    ***Isomerization***
Polymerization
   (photochem., photochem. and photophysics of liq. interfaces by second
   harmonic spectroscopy)
         ***nonlinear***
                           property
   (second-harmonic generation, photochem. and photophysics of liq.
   interfaces by second harmonic spectroscopy)
ANSWER 20 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
  ***Nonlinear***
                  optical properties of organic MIS structures
An important particularity of org. semiconductors is their strong coupling
between ***charge*** - ***transport*** and optical properties. By
applying a voltage on a Metal-Insulator-Semiconductor (MIS) structure
charges can be injected in the semiconductor under well-controlled
conditions. Based on these principles, new mechanisms of electrooptic
coupling and all-optical coupling (i.e., ***nonlinear*** optical)
phenomena appear in org. MIS devices. The use of these properties for the characterization of ***charge*** - ***transport*** and for
light-modulating devices are discussed. The example of the assocn. in a
MIS structure of electrooptic coupling and photo-cond. is discussed. It
leads to a new type of elec.-controlled ***photochromic*** device. A
review with 16 refs.
  ***nonlinear***
                  optical property org MIS review
Electrooptical effect
Optical ***nonlinear***
                           property
     ***nonlinear*** optical properties and electrooptical effect of
   org. MIS structures)
Optical instruments
                 ***nonlinear***
   (modulators,
                                   optical properties and electrooptical
   effect of org. MIS structures used in modulators)
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IT

L5

AB

IT

L5

AB

ΙT

ΙT

L5 TI

AB

ST

IT

IT

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L5
    ANSWER 21 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
    Novel second harmonic ***generation*** from intermolecular
TI
       ***charge*** -transfer complexes of styrylpyridinium salts in the
     crystalline state
ST
     styrylpyridinium salt charge transfer complex photophys; second harmonic
    generation styrylpyridinium ion pair; ***nonlinear*** optical property
     styrylpyridinium salt; photochem control second harmonic generation
       ***Photochromism***
IT
        (of intermol. charge-transfer complexes of styrylpyridinium salts in
        cryst. state)
IT
    Electron exchange and Charge transfer
        (photochem., novel second harmonic
                                            ***generation***
                                                               from intermol.
          ***charge*** -transfer complexes of styrylpyridinium salts in cryst.
        state)
              ***nonlinear***
IT
    Optical
                               property
        (second-harmonic generation, novel second harmonic ***generation***
                        ***charge*** -transfer complexes of styrylpyridinium
        from intermol.
        salts in cryst. state)
     140651-15-6, 1-Methyl-4-(4-nitrostyryl)pyridinium tetraphenylborate
IT
    RL: PEP (Physical, engineering or chemical process); PRP (Properties);
    PROC (Process)
        (novel second harmonic ***generation*** from intermol.
          ***charge*** -transfer complexes of styrylpyridinium salts in cryst.
     140651-16-7, 1-Methyl-4-(4-nitrostyryl)pyridinium methylbenzenesulfonate
    140651-17-8, 1-Methyl-4-(4-nitrostyryl)pyridinium perchlorate
    RL: PEP (Physical, engineering or chemical process); PRP (Properties);
     PROC (Process)
        (novel second harmonic
                               ***generation***
                                                   from intermol.
          ***charge*** -transfer complexes of styrylpyridinium salts in cryst.
        state in relation to)
    ANSWER 22 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
       . . were carried out in a guest/host system subjected to an external
     elec. field with and without the presence of a ***hole***
       ***transport*** agent. The guest was NNI, which has a moderate
     second-order ***nonlinear***
                                   optical coeff., the host was PMMA, and
          ***hole***
                        ***transport*** agent was DEH. In the presence of
    DEH, the lifetime of the cis ***isomer***
                                                of NNI detected by using the
    HGR technique is lengthened by the application of an elec. field of
    moderate strength.. . . obsd. when DEH is absent in the sample.
    result is interpreted as due to the stabilization of the cis
       ***isomer*** by the elec. field polarized DEH. The data obtained from
     SHG and EO measurements appear to corroborate the HGR result.
    Optical materials
        ( ***nonlinear*** , composite with PMMA, holog. gratings, laser
        induced relaxation of, elec. field effect on)
IT
     9011-14-7, PMMA
    RL: PRP (Properties)
        (contg. ***nonlinear*** optical compd. and
                                                        ***hole***
          ***transport*** agent, elec. field effect on laser-induced holog.
       grating relaxation of)
    ANSWER 23 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
AB
       . . were carried out in a guest/host system subject to an external
     elec. field with and without the presence of a    ***hole***
       ***transport*** agent. The guest is 1-(4-nitrophenylazo) 2-naphthyl
     isobutyrate (NNI), which has moderate second order ***nonlinear***
    optical ( ***NLO*** ) coeff., the host is PMMA and the
       ***transport*** agent is p-diethylaminobenzaldehyde diphenylhydrazone
     (DEH). In the presence of DEH, the lifetime of the cis ***isomer***
    of NNI detected by using the HGR technique is lengthened by the
    application of an elec. field of moderate strength.. . . not obsd. when
    DEH is absent in the sample. The result is interpreted as due to the
    stabilization of the cis- ***isomer*** by the elec. field polarized
    DEH. The data obtained from SHG and EO measurements appear to corroborate
    the HGR result.
    Electrooptical effect
    Optical ***nonlinear***
                              property
        (of photorefractive polymer, elec. field dependence of laser-induced
       holog. grating relaxation in relation to)
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ANSWER 24 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
ΤI
     Photorefractivity in doped ***nonlinear*** organic polymers
AB
     The properties of a new class of materials exhibiting the photorefractive
     effect, doped optical ***nonlinear*** org. polymers, are described.
     Photorefraction (at 647.1 nm) is established by a combination of hologram
     erasability, correlation with photocond. and electrooptic response, and
     enhancement by external fields in numerous samples (178-533 .mu.m thick)
            ***nonlinear***
                             epoxy materials doped with
                                                            ***hole***
     of 2
     ***transport*** agents based on p-(diethylamino)benzaldehyde diphenylhydrazone. Diffraction efficiencies .ltoreq.0.1% are obsd. at
     bias fields near 100 kV/cm. A useful property of these materials is that
                     ***nonlinear*** chromophores is partially reversible,
     poling of the
     permitting partial control of the grating readout independent of the
     space-charge field formed. The polarization anisotropy. . . than the
     index gratings, and (b) the phase shift of the index grating is near
     90.degree., which cannot occur via ***photochromism*** , heating, or
     any other process except photorefractivity.
                                                  epoxy resin; ***hole***
                              ***nonlinear***
ST
     photorefractivity doped
       ***transport***
                         agent epoxy photorefractivity
     Electrooptical effect
IT
        (of epoxy resins contg.
                                  ***hole***
                                                  ***transport***
                                                                    agents,
        photorefractivity in relation to)
IT
     Epoxy resins, properties
     RL: PRP (Properties)
                                ***nonlinear***
        (photorefractivity of
                                                   optical, doped with
                        ***transport***
          ***hole***
                                          agents)
     Optical materials
          ***nonlinear*** , epoxy resins doped with
                                                        ***hole***
          ***transport***
                           agents, photorefractivity of)
IT
     68189-23-1
                 83890-47-5
     RL: PRP (Properties)
          ***hole***
                          ***transport***
                                             agents,
                                                      ***nonlinear***
                                                                         optical
        epoxy resins doped with, photorefractivity of)
     125061-60-1
                  128611-17-6
     RL: PRP (Properties)
                                                   optical, doped with
        (photorefractivity of
                               ***nonlinear***
                         ***transport***
                                           agents)
L5
     ANSWER 25 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
TI
     Novel second harmonic
                             ***generation***
                                                from intermolecular
       ***charge*** -transfer complexes of styrylpyridinium tetraphenylborate
IT
       ***Photochromism***
        (of methyl(nitrostyryl)pyridinium tetraphenylborate)
IT
     Optical ***nonlinear*** property
        (harmonic
                    ***generation*** , second, from ***charge*** -transfer
        complexes of styrylpyridinium)
L5
     ANSWER 26 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
                 ***charge***
AΒ
     Defects and
                                    ***transport*** play crit. roles in the
     photorefractive behavior of electrooptic materials. Thus the
     photorefractive effect can be used as a noninvasive.
ΙT
       ***Photochromism***
        (of chromium-doped gallium arsenide)
IT
               ***nonlinear***
                               property
        (four-wave mixing, of chromium-doped gallium arsenide)
IT
     7440-47-3, Chromium, properties
     RL: PRP (Properties)
          ***nonlinear***
                             charge transfer and low temp. photorefractive
        effects in gallium arsenide contg.)
L5
     ANSWER 27 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
     Configuration of vinyl radicals. The generation and trapping of each
TI
     member of a configurationally ***isomeric*** pair of vinyl radicals
                       vinyl free radicals were generated by treating Na
AB
       ***Isomeric***
     naphthalenide with cis- and trans-3-chloro-3-hexene at 0.degree. and
     27.degree. using tetrahydrofuran and 1,2-dimethylethane as solvents.
     only detectable products were cis- and trans-3-hexene in 98% yield with
                ***isomer*** predominating. Since these results excluded
     the possibility of a single linear sp radical or 2 ***isomeric***
     configurationally stable, ***nonlinear*** sp2 vinyl free radicals, it was concluded that the initial ***charge*** -transfer reaction
       ***generates***
                            ***nonlinear*** vinyl radical which is capable of
                        a
```

L5

facile inversion at the radical site, but the radical is trapped before complete equilibration with its configurational \*\*\*isomer\*\*\* is achieved. A mechanism by which the interconverting free radical can be trapped is postulated.

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=> d all 4-27
    ANSWER 4 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
L5
AN
     2003:693438 CAPLUS
DN
     140:383024
ED
     Entered STN: 05 Sep 2003
    Holographic gratings in hybrid sol-gel films
TI
    Raschella, R.; Marino, I.-G.; Lottici, P. P.; Bersani, D.; Lorenzi, A.;
ΑU
    Montenero, A.
     INFM, Univ. degli Studi di Parma, Parma, 43100, Italy
CS
SO
     Proceedings of SPIE-The International Society for Optical Engineering
     (2003), 5123 (Advanced Optical Devices, Technologies, and Medical
     Applications), 117-124
     CODEN: PSISDG; ISSN: 0277-786X
     SPIE-The International Society for Optical Engineering
PB
     Journal
DT
LA
     English
     74-8 (Radiation Chemistry, Photochemistry, and Photographic and Other
CC
     Reprographic Processes)
    Org.-inorg. films based on SiO2, contg. Disperse Red 1 (DR1),
AB
       ***carbazole***
                        units and 2,4,7-trinitro-9-fluorenone (TNF), have been
     prepd. by a sol-gel technique. Diffraction gratings have been produced
     through different effects: photoinduced birefringence and
     photorefractivity, using 488.0 nm and 632.8 nm light, resp.
    polarization holog. birefringence gratings have been investigated and
     diffraction efficiencies higher than those obtained by light intensity
     modulation have been measured. The study of the temporal behavior of the
     diffraction efficiencies makes possible the identification of different
    processes involved in the DR1 mol. orientation: angular hole burning
     (AHB), angular redistribution (AR), Cis mols. gain has been detd. by
     two-beam coupling (2BC) measurements. The effect of the polarization of
     the writing beams on the grating and of a circularly polarized
       ***photoisomerizing***
                              radiation during grating erasure has been
     interpreted in terms of an orientation contribution to the grating
     polarization holog birefringence grating hybrid sol gel film
st
IT
       ***Isomerization***
        (cis-trans, photochem.; recording of holog. gratings in hybrid
        org.-inorg. films based on SiO2 sol-gel films contg. Disperse Red 1 and
          ***carbazole***
                           units and trinitrofluorenone)
    Birefringence
        (photoinduced; recording of holog. gratings in hybrid org.-inorg. films
       based on SiO2 sol-gel films contg. Disperse Red 1 and ***carbazole***
        units and trinitrofluorenone)
    Holographic diffraction gratings
    Holographic recording materials
     Hybrid organic-inorganic materials
     Molecular orientation
     Photorefractive effect
     Sol-gel processing
        (recording of holog, gratings in hybrid org.-inorg, films based on SiO2
        sol-gel films contg. Disperse Red 1 and
                                                  ***carbazole***
                                                                    units and
       trinitrofluorenone)
       ***Nonlinear***
                        optical properties
        (two-beam-coupling; recording of holog. gratings in hybrid org.-inorg.
        films based on SiO2 sol-gel films contq. Disperse Red 1 and
          ***carbazole***
                           units and trinitrofluorenone)
     7647-01-0, Hydrochloric acid, uses
IT
    RL: CAT (Catalyst use); USES (Uses)
        (recording of holog. gratings in hybrid org.-inorg. films based on SiO2
        sol-gel films contg. Disperse Red 1 and ***carbazole***
                                                                    units and
        trinitrofluorenone)
IT
     685091-04-7
    RL: PEP (Physical, engineering or chemical process); PYP (Physical
    process); PROC (Process)
```

(recording of holog. gratings in hybrid org.-inorg. films based on SiO2

```
sol-gel films contg. Disperse Red 1 and
                                                 ***carbazole***
                                                                    units and
        trinitrofluorenone)
IT
     919-30-2, 3-Aminopropyltriethoxysilane
                                              2872-52-8, Disperse Red 1
     24801-88-5, 3-(Triethoxysilyl)propylisocyanate
                                                     73500-82-0,
       ***Carbazole***
                       -9-carbonyl chloride
     RL: PEP (Physical, engineering or chemical process); PYP (Physical
     process); RCT (Reactant); PROC (Process); RACT (Reactant or reagent)
        (recording of holog. gratings in hybrid org.-inorg. films based on SiO2
        sol-gel films contg. Disperse Red 1 and
                                                  ***carbazole***
        trinitrofluorenone)
IT
     129-79-3, 2,4,7-Trinitro-9-fluorenone
     RL: PEP (Physical, engineering or chemical process); PYP (Physical
     process); PROC (Process)
        (sensitizer; recording of holog. gratings in hybrid org.-inorg. films
        based on SiO2 sol-gel films contg. Disperse Red 1 and
                                                                ***carbazole***
        units and trinitrofluorenone)
              THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD
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RE
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L_5
     ANSWER 5 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
     2003:99139 CAPLUS
AN
DN
     138:375505
ED
     Entered STN: 09 Feb 2003
     Molecular engineering of dye-doped polymers for optoelectronics
ΤI
ΑU
     Ishchenko, Alexander
     Institute of Organic Chemistry, National Academy of Sciences of Ukraine,
CS
     Kiev, 02094/94, Ukraine
SO
     Polymers for Advanced Technologies (2002), 13(10-12), 744-752
     CODEN: PADTE5; ISSN: 1042-7147
PB
     John Wiley & Sons Ltd.
DT
     Journal
LA
     English
CC
     73-4 (Optical, Electron, and Mass Spectroscopy and Other Related
     Properties)
     The main approaches to mol. engineering of prospective dye-doped polymer
     matrixes for optoelectronics were considered. The advantages of such
     matrixes over polymers and dyes sep. were analyzed. These advantages can
     be realized in the case where a doping of polymers by low-mol.-wt. dyed
     impurities does not essentially distort spectral, luminescent and
       ***nonlinear***
                         optical properties of the individual components of
     polymer compn. The influence of polymer nature and chem. constitution of
     org. dyes on photophys. and photochem. properties of these matrixes was
     analyzed. Processes of dye aggregation in polymers were characterized.
     Their influence on photophys. properties and the photochem. stability of
     dye-doped materials is discussed. The different approaches for struggle
     with it is offered. The influence of the method of introduction of a dye
     into a polymer on the output parameters of dyed materials was considered.
     The main paths of energy degrdn. of electronic excitation in such
     materials - internal conversion, electron transfer and
       ***photoisomerization***
                                  - were analyzed. The role of dyes in the
     process of generation, in recombination and in the carriage of charges in
     photoconductive polymers is discussed. The prospects for the application
     of dye-doped polymer materials as passive Q-switches of solid-state
     lasers, as active laser media with large Stokes shift, as luminescent
     solar converters and as electroluminescent emitters were demonstrated.
     dye doped conjugated polymer optoelectronics polyurethane
     electroluminescent device; solid state laser solar converter dye doped
```

```
conjugated polymer
IT
     Polymers, properties
     RL: DEV (Device component use); PRP (Properties); USES (Uses)
        (conjugated; mol. engineering of dye-doped polymers for
        optoelectronics)
IT
     Doping
    Dyes
     Electroluminescent devices
     Optoelectronics
     Solid state lasers
        (mol. engineering of dye-doped polymers for optoelectronics)
IT
     Polyurethanes, properties
     RL: DEV (Device component use); PRP (Properties); USES (Uses)
        (mol. engineering of dye-doped polymers for optoelectronics)
IT
     51829-02-8
                  80993-83-5
                               101491-20-7
     RL: DEV (Device component use); MOA (Modifier or additive use); PRP
     (Properties); USES (Uses)
        (mol. engineering of dye-doped polymers for optoelectronics)
IT
     25067-59-8,
                 ***Polyvinylcarbazole***
                                                 55774-96-4,
     Poly-N-epoxypropylcarbazole
     RL: DEV (Device component use); PRP (Properties); USES (Uses)
        (mol. engineering of dye-doped polymers for optoelectronics)
              THERE ARE 49 CITED REFERENCES AVAILABLE FOR THIS RECORD
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AN
DN
     139:187903
ED
     Entered STN: 12 Jan 2003
ΤI
     CdSe/CdS core/shell quantum dots as sensitizer of a photorefractive
    polymer composite
     Binks, D. J.; Bant, S. P.; West, D. P.; O'Brien, P.; Malik, M. A.
ΑU
     Dept. Chem., Univ. Manchester, Manchester, UK
CS
     Journal of Modern Optics (2003), 50(2), 299-310
SO
     CODEN: JMOPEW; ISSN: 0950-0340
    Taylor & Francis Ltd.
PΒ
     Journal
DT
LA
     English
     73-11 (Optical, Electron, and Mass Spectroscopy and Other Related
CC
     Properties)
     Section cross-reference(s): 74
     CdSe/CdS core/shell, tri-n-octylphosphine oxide passivated, quantum dots
AB
    are used to sensitize a photorefractive polymer composite. The composite
    also consists of poly(N- ***vinylcarbazole*** ) as the nominally
                        ***transporting***
       ***charge***
                                            matrix and an electrooptic
     chromophore. The efficacy of sensitization and consequent photorefractive
     performance is investigated using transmission spectroscopy and
     ellipsometry, two-beam coupling and degenerate four-wave mixing expts.
     The photorefractive nature of the photoinduced grating is confirmed by the
     observation of asym. two-beam coupling. Four-wave mixing reveals record
     diffraction efficiencies for a nanoparticle-sensitized photorefractive
     polymer at the field levels applied (1.3% at 70 V. .mu.m-1). A recently
     developed anal. technique is used to ext. space-charge field rise time
     values from degenerate four-wave mixing transients. In turn, anal. of the
     dependence of the rise time on applied field is used to det. the
     zero-field charge dissocn. efficiency to be 3.6.times. 10-5.+-.0.5.times.
     10-5. It is further shown that the magnitude of this parameter accounts
     for most of the difference in photorefractive response rate between the
     present material and a similar C60 sensitized composite.
     photorefractive polymer composite cadmium sulfide selenide quantum dot
     sensitizer
       ***Isomerization***
IT
        (cis-trans, photochem.; photorefractive poly( ***vinylcarbazole***
        )-based composite contq. (ethylhexyloxydimethylnitrophenylazo)benzene
        sensitized by octylphosphine oxide passivated CdSe/CdS quantum dots in
        relation to)
     Degenerate four wave mixing
     Ellipsometry
     Holographic diffraction gratings
     Nanoparticles
     Photorefractive effect
     Photorefractive materials
     Quantum dot devices
     Refractive index
     Space charge
        (photorefractive poly( ***vinylcarbazole*** )-based composite contg.
        (ethylhexyloxydimethylnitrophenylazo)benzene sensitized by
       octylphosphine oxide passivated CdSe/CdS quantum dots)
IT
    Holographic recording materials
        (photorefractive poly( ***vinylcarbazole*** )-based composite contq.
        (ethylhexyloxydimethylnitrophenylazo)benzene sensitized by
        octylphosphine oxide passivated CdSe/CdS quantum dots in relation to)
IT
     Birefringence
        (transient; photorefractive poly( ***vinylcarbazole*** )-based
        composite contg. (ethylhexyloxydimethylnitrophenylazo)benzene
        sensitized by octylphosphine oxide passivated CdSe/CdS quantum dots)
       ***Nonlinear***
                        optical properties
        (two-beam-coupling; photorefractive poly( ***vinylcarbazole***
        )-based composite contg. (ethylhexyloxydimethylnitrophenylazo)benzene
        sensitized by octylphosphine oxide passivated CdSe/CdS quantum dots)
```

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78-50-2, Tri-n-octylphosphine oxide
                                           1306-23-6, Cadmium sulfide,
IT
     properties
                  1306-24-7, Cadmium selenide, properties
                                                           25067-59-8, Poly(N-
       ***vinylcarbazole*** )
                                 176681-76-8, EHDNPB
     RL: PEP (Physical, engineering or chemical process); PRP (Properties); PYP
     (Physical process); PROC (Process)
        (photorefractive poly( ***vinylcarbazole*** )-based composite contq.
        (ethylhexyloxydimethylnitrophenylazo) benzene sensitized by
        octylphosphine oxide passivated CdSe/CdS quantum dots)
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ΑN
     2003:12199 CAPLUS
DN
     138:278013
ED
     Entered STN: 07 Jan 2003
TI
     Optical nonlinearity of fullerene-doped polymer nanocomposites
ΑU
     Antipov, O. L.; Yurasova, I. V.; Domrachev, G. A.
CS
     Institute of Applied Physics, Russian Academy of Sciences, Nizhnii
     Novgorod, 603600, Russia
so
     Quantum Electronics (2002), 32(9), 776-780
     CODEN: QUELEZ; ISSN: 1063-7818
PB
     Turpion Ltd.
DT
     Journal
LA
     English
CC
     73-10 (Optical, Electron, and Mass Spectroscopy and Other Related
AΒ
     The mechanism of optical nonlinearity of new polymer composites based on a
     conducting polymer [poly(9- ***vinylcarbazole*** )] and fullerenes C70
                                                      ***nonlinear***
     and C60 was studied both exptl. and theor. The
     -optical studies of self-action and coupling of two 633-nm beams from a
     He-Ne laser were performed, and variations in the absorption spectra of
     the composites illuminated by a laser beam were studied. A giant inertial
     nonlinearity of org. materials is caused by the difference in
     polarizabilities of fullerene mols. and their anion radicals, which are
     formed upon absorption of photons and charge transfer by poly(9-
       ***vinylcarbazole*** ) mols.
                           ***carbazole***
ST
     fullerene polyvinyl
                                               ***nonlinear***
     property nanocomposite
IT
     Conducting polymers
     Nanocomposites
         ***Nonlinear***
                          optical susceptibility
         ***Photochromism***
     Photorefractive effect
        (optical nonlinearity of fullerene-doped polymer nanocomposites)
IT
     25067-59-8, Poly(9- ***vinylcarbazole*** ) 99685-96-8, Fullerene (C60)
     115383-22-7, Fullerene C70
     RL: PRP (Properties)
        (optical nonlinearity of fullerene-doped polymer nanocomposites)
RE.CNT
              THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE
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```
***photochromism***
        (prepn., properties and
                                                        of charge transfer
        compds. with Keggin anions and 2,2'-biquinolinium)
ΪT
     UV and visible spectra
        (solid-state and soln.; of charge transfer compds. with Keggin anions
        and 2,2'-biquinolinium)
     119-91-5, 2,2'-Biquinoline
                                  11089-20-6
                                               12027-43-9
                                                             51429-74-4
IT
     395071-70-2
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (for prepn. of charge transfer compds. with Keggin anions and
        2,2'-biquinolinium)
IT
     503014-03-7P
    RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and UV-visible spectra)
                    503014-13-9P
IT
     503014-11-7P
    RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of charge transfer compds. with Keggin anions and
        2,2'-biquinolinium)
IT
     503014-07-1P
                    503014-09-3P
                                   503014-15-1P
                                                  503014-17-3P
    RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
        (prepn., UV-visible spectra and ***photochromism*** )
IT
     503014-05-9P
    RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
        (prepn., UV-visible spectra and second-harmonic generation)
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grating as well as photorefractive grating were obsd. in the composite
    consisting of 1-n-butoxyl-2,5- dimethyl-4-(4'-nitrophenylazo) benzene
     (BDMNPAB): poly(N- ***vinylcarbazole*** ) (PVK): 2,4,7-trinitro-9-
    fluorenone (TNF) in a wt. ratio of 44:55:1. Based on the quick
    translation technique of two-beam coupling exptl. geometry, the authors
    measured the electroabsorption grating to be 6-7\ \text{cm-1}, and
                             effect also brought an absorption grating of 2-3
       ***photoisomeric***
              ***Photoisomeric***
                                    effect caused index grating was sepd. from
    the others by analyzing the dynamic behavior of the gratings, and an
    amplitude between 1 .times. 10-4.apprx.2 .times. 10-4 was obtained.
                                                                 chromophore
    multi grating photorefractive composite
                                               ***nonlinear***
                                        ***photoisomerization***
    azo dye; photorefractive grating
    electroabsorption two beam coupling
       ***Nonlinear***
                        optical properties
        (beam coupling; multi-grating in photorefractive composites contg.
          ***nonlinear***
                           chromophore azo-dye 1-n-butoxyl-2,5-dimethyl-4-(4'-
       nitrophenylazo)benzene)
    Azo dyes
    Electric field effects
    Electrooptical absorption
    Photorefractive gratings
        (multi-grating in photorefractive composites contg.
                                                              ***nonlinear***
       chromophore azo-dye 1-n-butoxyl-2,5-dimethyl-4-(4'-
       nitrophenylazo) benzene)
       ***Isomerization***
        ( ***photoisomerization*** ; multi-grating in photorefractive
                            ***nonlinear***
       composites contg.
                                              chromophore azo-dye
        1-n-butoxyl-2,5-dimethyl-4-(4'-nitrophenylazo)benzene)
                                             25067-59-8, Poly(N-
     129-79-3, 2,4,7-Trinitro-9-fluorenone
       ***vinylcarbazole*** ) ~ 210281-60-0
    RL: DEV (Device component use); PRP (Properties); USES (Uses)
                                                              ***nonlinear***
        (multi-grating in photorefractive composites contg.
        chromophore azo-dye 1-n-butoxyl-2,5-dimethyl-4-(4'-
      nitrophenylazo)benzene)
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    ANSWER 11 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
     1998:429870 CAPLUS
     129:181774
    Entered STN: 13 Jul 1998
       ***Photoisomeric*** -chromophore-doped polymers for directional spatial
     filtering by optical threshold processing
    Egami, C.; Suzuki, Y.; Aoshima, Y.; Sugihara, O.; Okamoto, N.
    Faculty of Engineering, Department of Electrical and Electronic
    Engineering, Shizuoka University, 3-5-1, Johoku, Hamamatsu, Shizuoka, 432,
    Journal of the Optical Society of America B: Optical Physics (1998),
     15(7), 1985-1991
    CODEN: JOBPDE; ISSN: 0740-3224
    Optical Society of America
    Journal
    English
    73-10 (Optical, Electron, and Mass Spectroscopy and Other Related
    Properties)
    Section cross-reference(s): 74
    We demonstrate spatial filtering by using the azo-chromophore-based
```

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optical effect of self-beam polarization modulation and
       ***nonlinear***
     optical threshold processing. The real-time image-processing scheme is
     described by use of an optical configuration composed of an
     action-beam-probe-beam system, Fourier-transform operation, and an
     azo-chromophore-doped film as a spatial filter. As Disperse Red 1 dye (DR1) doped in an N-poly( ***vinylcarbazole*** ) (PVK) film with
     N-ethylcarbazole (ECZ) of low glass-transition temp. exhibits
     self-polarization rotation after thin-film transmission, a linearly
     polarized probe beam in the film is obsd., even through a crossed
     analyzer. In addn., this film permits optical threshold processing by
     introducing an action beam with different intensity and wavelength from
     those of the probe beam with object information. Changing the action-beam
     intensity results in a decrease or an increase in the transmitted
     probe-beam power through the analyzer. When a DR1-PVK-ECZ film is placed
     at the Fourier plane on the probe-beam path, the filter functions can be
     modulated in real time by manipulation of the intensity ratio between the
     action beam and the spatial-frequency components at the film plane. The
     spatial frequencies of an original input image can be manipulated as they
     pass through the film. These effects are responsible for the
                                 of DR1 mols. in the low-glass-transition-temp.
       ***photoisomerization***
     polymer matrix.
       ***photoisomeric***
                             chromophore doped polymer; directional spatial
     filtering optical threshold processing
       ***Isomerization***
                                  ***photoisomeric*** -chromophore-doped
        (cis-trans, photochem.;
        polymers for directional spatial filtering by optical threshold
        processing)
     Memory devices
         ***Nonlinear***
                           optical materials
     Optical films
     Optical filters
     Optical modulation
     Spatial light modulators
           ***photoisomeric***
                                -chromophore-doped polymers for directional
        spatial filtering by optical threshold processing)
     86-28-2, N-Ethylcarbazole 2872-52-8, Disperse Red 1
                                                              25067-59-8,
                  ***carbazole***
     Poly(vinyl
     RL: PRP (Properties); TEM (Technical or engineered material use); USES
          ***photoisomeric*** -chromophore-doped polymers for directional
        (
        spatial filtering by optical threshold processing)
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ΑN
DN
     129:142426
ED
     Entered STN: 09 Jul 1998 ·
TI
     Development of self adaptive image processing system by use of a
       ***photoisomeric***
                            organic dye film
     Egami, Chikara; Sugihara, Okihiro; Fujimura, Hisashi; Okamoto, Naomichi
ΑU
     Fac. Eng., Shizuoka Univ., Hamamatsu, Japan
CS
     Shizuoka Daigaku Kogakubu Kenkyu Hokoku (1997), 48, 1-6
SO
     CODEN: SDKKAT; ISSN: 0583-0915
PB
     Shizuoka Daigaku Kogakubu
     Journal; General Review
DΤ
LA
     Japanese,
     74-0 (Radiation Chemistry, Photochemistry, and Photographic and Other
CC
     Reprographic Processes)
     Section cross-reference(s): 73
     A review with 8 refs. We demonstrate spatial filtering using the
AB
     azo-chromophore-based
                            ***nonlinear*** optical effect of the self-beam
     polarization-modulation. The real-time image processing scheme is
     described using the optical configuration, composed of an action-probe
     beam system. Fourier-transform operation and an azo-chromophore-doped
     film as a spatial filter. As Disperse Red 1 doped in the N-
       ***polyvinylcarbazole***
                                  film with N-ethylcarbazole, of low
     glass-transition-temp., exhibits the self polarization-modulation after
     the thin film transmission, a linearly-polarized probe beam into the film
     is obsd. even through a crossed analyzer. When the DR1/PVK/ECZ film is
     placed at the Fourier plane on the probe beam path, the filter functions
     can be modulated in real time. The spatial frequencies of an original
     input image can be manipulated as they pass through the film. This effect
     is responsible for the ***photoisomerization***
                                                         of DR1 mols. in the
     low-glass-transition-temp. polymer matrix.
ST
     review self adaptive image processing;
                                              ***photoisomeric***
     film review
IT
     Imaging
     Optical films
     Optical filters
     Spatial light modulators
        (development of self adaptive image processing system by use of a
          ***photoisomeric***
                               org. dye film)
IT
     Memory effect
        (optical; development of self adaptive image processing system by use
               ***photoisomeric*** org. dye film)
       ***Isomerization***
           ***photoisomerization*** ; development of self adaptive image
       processing system by use of a
                                        ***photoisomeric***
                                                              org. dye film)
                               2872-52-8, C.I. Disperse Red 1
     86-28-2, N-Ethylcarbazole
                                                                  25067-59-8,
     N- ***Polyvinylcarbazole***
     RL: PRP (Properties); TEM (Technical or engineered material use); USES
        (development of self adaptive image processing system by use of a
          ***photoisomeric***
                               org. dye film)
     ANSWER 13 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
L5
     1998:261567 CAPLUS
ΑN
DN
     128:328463
ED
     Entered STN: 08 May 1998
     Organic-inorganic solids by sol-gel processing: optical applications
TI
     Boilot, J. -P.; Biteau, J.; Chaput, F.; Gacoin, T.; Brun, A.; Darracq, B.;
     Georges, P.; Levy, Y.
     Groupe de Chimie du Solide, Physique de la Matiere Condensee, URA CNRS
     1254 D, Ecole Polytechnique, Palaiseau, 91128, Fr.
SO
     Pure and Applied Optics (1998), 7(2), 169-177
     CODEN: PAOAE3; ISSN: 0963-9659
PΒ
     Institute of Physics Publishing
DT
     Journal
LΑ
     English
     73-10 (Optical, Electron, and Mass Spectroscopy and Other Related
     Properties)
     Section cross-reference(s): 38, 66
AB
     We have developed pure inorg. and hybrid org.-inorg. solids by sol-gel
     processing of silicon alkoxides. Dense gels were obtained at room temp.,
     in any desired shape, including thin films, and we are able to trap org.
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mols. or clusters. Gels can be polished and exhibit optical transmission
     in the visible range similar to that of high-temp. silica glasses.
    Concerning gel-mol. and gel-nanocrystal composites, the diversity of the
    organofunctional alkoxide precursors allows us to modify the optical
    properties by changing the nature and the strength of chem. and electronic
    interactions between the optically active guest system and the solid host
    matrix.
    sol gel org inorg optical properties; xerogel pyrromethene optical
    pumping; merocyanine spiro oxazine
                                         ***photochromic***
                                                              sol qel; laser
    written grating DR1 sol gel; photorefractive sol gel ***carbazole***
       ***Photochromism***
        (and reverse- ***photochromism***
                                             of spiro-oxazine in sol-gel
       matrix)
    Solid state lasers
    Xerogels
        (dense xerogels for tunable solid state laser applications)
    Laser induced grating
        (in disperse red 1 doped sol-gel matrix)
    Photorefractive effect
        (in sol-gel materials using functionalized silicon alkoxides)
    Optical pumping
        (of pyrromethene-597 in dense xerogel)
    Optical properties -
    Optical transmission
    Sol-gel processing
        (optical applications of org.-inorg. solids by sol-gel processing)
       ***Nonlinear*** optical properties
        (two-beam coupling; in photorefractive sol-gel materials)
     137829-79-9, Pyrromethene-597
    RL: DEV (Device component use); MOA (Modifier or additive use); USES
     (Uses) .
        (dense xerogels for tunable solid state laser applications)
     2872-52-8, Disperse red 1
    RL: DEV (Device component use); MOA (Modifier or additive use); USES
     (Uses)
        (laser-written gratings in doped sol-gel matrix)
     206867-72-3D, derivs. 206867-73-4D, derivs.
     RL: PEP (Physical, engineering or chemical process); PRP (Properties);
     PROC (Process)
           ***photochromism***
                                 and reverse- ***photochromism***
                                                                     in sol-qel
       matrix)
    78-08-0, Vinyl triethoxysilane
                                    998-30-1, Triethoxysilane
    Methyltriethoxysilane
    RL: PEP (Physical, engineering or chemical process); PROC (Process)
           ***photochromism*** and reverse- ***photochromism***
        spiro-oxazine in sol-gel matrix from)
                ***Carbazole***
                                 , silicon alkoxide deriv.
    Disperse red 1, silicon alkoxide deriv.
    RL: PEP (Physical, engineering or chemical process); PRP (Properties);
     PROC (Process)
        (photorefractive sol-gel materials using)
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     1997:703445 CAPLUS
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     128:8701
     Entered STN: 08 Nov 1997
ED
    New photorefractive polymers for real-time holography
ΤI
     Schloter, S.; Hofmann, U.; Hagen, R.; Hohle, C.; Ewert, K.; Strohriegl,
AU
     P.; Eisenbach, C.-D.; Schmidt, H.-W.; Haarer, D.
    Lehrstuhl Experimentalphysik IV, Universitaet Bayreuth, Bayreuth, 95440,
CS
SO
     Proceedings of SPIE-The International Society for Optical Engineering
     (1997), 3144 (Xerographic Photoreceptors and Organic Photorefractive
     Materials II), 142-153
     CODEN: PSISDG; ISSN: 0277-786X
PB
     SPIE-The International Society for Optical Engineering
DT
     Journal
LΑ
     English
     74-8 (Radiation Chemistry, Photochemistry, and Photographic and Other
     Reprographic Processes)
     Section cross-reference(s): 38
AB
     We investigated polymeric materials based on polysiloxane (PSX),
     polymethylmethacrylate (PMMA), polyurethane (PU), as well as a
     triphenylamine-based glass (DRDCTA) with respect to their photorefractive
     properties. Elec.-field dependencies of the two-beam coupling gain,
     diffraction efficiencies, refractive index amplitudes and holog.
     rise-times could be obtained by means of two-wave mixing and degenerate
     four-wave mixing measurements. The examd. PSX polymer systems were
     composed of a photoconducting polysiloxane host doped with
     trinitrofluorenone (TNF) as a sensitizing moiety and various chromophores,
     namely, an azo deriv., a stilbene deriv. and a tolan deriv. Due to their
     comparatively low glass-transition temps. Tg, an orientational enhancement
     of the photorefractive properties was obsd. Furthermore, the influence of
       ***photoisomerization***
                                 (based on trans-cis-trans cycles) on the holog.
     properties could be detd. for the different chromophores. In addn., a
     class of fully functionalized polymers with azo chromophores and
       ***carbazole*** -units covalently attached to PMMA- and PU-backbones was
     synthesized. These systems show comparatively high glass transition
     temps. of more than 80.degree.C. The third type of materials investigated
                                                  ***carbazole***
     is a glass of triphenylamine with attached
                 -chromophore moieties. It has a glass transition temp. of
     120.degree.C. For the high-Tg materials, poling procedures - essential
     for the photorefractive properties - could be monitored in-situ by
     second-harmonic generation. Abs. values for the
                                                       ***nonlinear***
     Pockels coeffs. .chi.(2)(-.omega.;.omega.,0) have been obtained by
     electro-optical measurements.
     photorefractive polymer real time holog diffraction; photoconducting
ST
     polysiloxane host trinitrofluorenone dopant holog
IT
     Testing of materials
        (nondestructive; photorefractive polymers for real-time holog.)
       ***Isomerization***
           ***photoisomerization*** ; photorefractive polymers for real-time
        holog.)
     Degenerate four wave mixing
     Dielectric polarization
     Holography
     Optical diffraction
     Photorefractive materials
     Refractive index
     Second-harmonic generation
```

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Two wave mixing
        (photorefractive polymers for real-time holog.)
     Polymers, properties
     Polysiloxanes, properties
     RL: PRP (Properties); TEM (Technical or engineered material use); USES
     (Uses)
        (photorefractive polymers for real-time holog.)
     Polyurethanes, properties
IT
     Polyurethanes, properties
     RL: PRP (Properties); TEM (Technical or engineered material use); USES
     (Uses)
        (polyamine-; photorefractive polymers for real-time holog.)
IT
     Polyamines
     Polyamines
     RL: PRP (Properties); TEM (Technical or engineered material use); USES
     (Uses)
        (polyurethane-; photorefractive polymers for real-time holog.)
                             198827-68-8
IT
     129-79-3
                153800-56-7
                                            198827-69-9
     RL: MOA (Modifier or additive use); USES (Uses)
        (photorefractive polymers for real-time holog.)
IT
     9011-14-7, Polymethylmethacrylate
                                        150528-92-0
                                                        181427-94-1
     198827-72-4
                   198827-73-5
                                 198827-75-7
                                               198827-84-8
     RL: PRP (Properties); TEM (Technical or engineered material use); USES
     (Uses)
        (photorefractive polymers for real-time holog.)
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AN
DN
     127:294024
ED
     Entered STN: 24 Oct 1997
                               ***nonlinear***
                                                 optical properties of novel
TI
     Linear and second order
       ***photochromic***
                            materials
     Zinger, Baruch; Schaer, Pnina; Berkovic, Gary; Meshulam, Guilia; Kotler,
AU
     Zvi; Shapiro, Lev; Mazor, Royi; Khodorkovsky, Vladimir
CS
     Nonlinear Optics Group SOREQ NRC, Yavne, 81800, Israel
SO
     Proceedings of SPIE-The International Society for Optical Engineering
     (1997), 3135 (Precision Plastic Optics for Optical Storage, Displays,
     Imaging, and Communications), 71-78
     CODEN: PSISDG; ISSN: 0277-786X
PB
     SPIE-The International Society for Optical Engineering
DT
    Journal
LA
    English
CC
     37-5 (Plastics Manufacture and Processing)
     Section cross-reference(s): 36, 73
AB
     A new unique
                    ***photochromic***
                                         material which is based on a
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reversible formation - cleavage of a C-C bond is described. The solution bindon deriv. (.lambda.max = 480 nm in toluene) undergoes
                                                                   The bicyclic
    photochem. and/or thermally induced ring opening to form the
       ***isomer***
                     (.lambda.max= 640 nm in toluene). This
     form presents a conjugated donor-acceptor system and exhibits a
    considerable second-order optical nonlinearity as found by field induced
    SHG (EFISH) measurements.
                                  ***Photochromic*** conversion was also
    obsd. in the cryst. form indicated visually by a crystal red-to-green
    color change. The reversible ring opening - closure process in liq. and
    polymeric solns. [PMMA and poly( ***vinylcarbazole*** )] was studied.
                                             ***NLO***
    Optical and thermal switching and the
                                                         efficiency of the
    guest-host polymer system are described.
                                             ***isomer***
    bindon dimethylaminocinnamic aldehyde
       ***photochromic***
                           conversion;
                                          ***nonlinear***
                                                            optical property
    biindantrione dimethylaminocinnamic aldehyde; solvatochromism
       ***photochromism***
                             indan deriv polymer host; polymethylmethacrylate
                             ***NLO*** ;
    host indantrione deriv
                                             ***polyvinylcarbazole***
    indantrione deriv
                        ***NLO***
       ***Isomerization***
        (cis-trans, photochem. and thermally induced; linear and second order
          ***nonlinear*** optical properties of biindantrione deriv.
          ***photochromic***
                              material in polymer hosts)
    Optical switching
         ***Photochromic***
                              materials
         ***Photochromism***
                    ***nonlinear***
                                      optical properties
    Second-order
    Solvatochromism
                                                     optical properties of
        (linear and second order
                                  ***nonlinear***
                              ***photochromic*** material in polymer hosts)
        biindantrione deriv.
    56-23-5, Carbon tetrachloride, uses 67-66-3, Chloroform, uses
                                           75-05-8, Acetonitrile, uses
                  71-43-2, Benzene, uses
    DMSO, uses
    75-09-2, uses 108-88-3, Toluene, uses
                                               110-82-7, Cyclohexane, uses
     123-91-1, Dioxane, uses
                               9011-14-7, PMMA
                                                 25067-59-8, Poly(
       ***vinylcarbazole*** )
    RL: NUU (Other use, unclassified); USES (Uses)
        (linear and second order ***nonlinear***
                                                     optical properties of
                               ***photochromic***
        biindantrione deriv.
                                                    material in polymer hosts)
    197023-27-1P
                    197023-28-2P
    RL: PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); PREP
     (Preparation); RACT (Reactant or reagent)
                                   ***nonlinear***
        (linear and second order
                                                     optical properties of
                              ***photochromic***
                                                    material in polymer hosts)
       biindantrione deriv.
                       6203-18-5, p-(Dimethylamino)cinnamic aldehyde
    1707-95-5, Bindon
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (linear and second order ***nonlinear***
                                                     optical properties of
                              ***photochromic***
                                                    material in polymer hosts)
        biindantrione deriv.
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    ANSWER 16 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
    1997:579455 CAPLUS
    127:270409
    Entered STN: 11 Sep 1997
       ***Carbazole*** -based multifunctional molecules for photorefractive
    applications
    Wang, Qing; Gharavi, Alireza; Li, Wenjie; Yu, Luping
    Department of Chemistry and James Frank Institute, The University of
    Chicago, Chicago, IL, 60637, USA
    Polymer Preprints (American Chemical Society, Division of Polymer
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Chemistry) (1997), 38(2), 516-517
     CODEN: ACPPAY; ISSN: 0032-3934
PB
     American Chemical Society, Division of Polymer Chemistry
DT
     Journal
LA
     English
     74-8 (Radiation Chemistry, Photochemistry, and Photographic and Other
CC
     Reprographic Processes)
     A multifunctional
                         ***photochromic***
                                              mols. consist of a
AB
       ***carbazole***
                         part with a
                                     ***nonlinear***. optical chromophore unit
     attached via an alkyl link to the N atom of the ***carbazole***
     These compds. exhibit dual functions: photocond. and a second order
                         optical activity. The photocharge transport and the
       ***nonlinear***
     electrooptic functions are performed by two sep. constituents.
     Photorefractive effect of ethyl(hexylsulfonylphenylazophenyl)aminododecylc
     arbazole (I) was studied along with dynamic holog. recording in a film
     contg. I and 0.9% of 2,4,7-trinitro-9-fluorenone.
                     ***carbazole***
                                        based multifunctional mol holog;
ST
     photorefractive
       ***nonlinear*** optical substituent ***carbazole***
     photorefractive
IT
     Holography
     Photorefractive effect
     Photorefractive materials
        (photorefractive effect and dynamic holog, recording using
          ***carbazole*** -based multifunctional mols.)
       ***Nonlinear*** optical properties
IT
        (two beam-coupling; of photorefractive ***carbazole*** -based
        multifunctional mols.)
IT
     196201-41-9
     RL: PRP (Properties); TEM (Technical or engineered material use); USES
     (Uses)
        (photorefractive effect and dynamic holog. recording using
          ***carbazole*** -based multifunctional mols.)
IT
     129-79-3, 2,4,7-Trinitro-9-fluorenone
     RL: TEM (Technical or engineered material use); USES (Uses)
        (photorefractive effect and dynamic holog. recording using
          ***carbazole*** -based multifunctional mols.)
     ANSWER 17 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
L5
     1997:501363 CAPLUS
AN
DN
     127:211660
     Entered STN: 08 Aug 1997
ED
     Transient absorption spectroscopy of semiconductor nanoclusters
TI
ΑU
     Kamat, Prashant V.
     Radiation Lab., Univ. Notre Dame, Notre Dame, IN, 46556, USA
CS
SO
     Reza Kenkyu (1997), 25(6), 417-424
     CODEN: REKEDA; ISSN: 0387-0200
PB
     Reza Gakkai
     Journal; General Review
DT.
LA
     English
     73-0 (Optical, Electron, and Mass Spectroscopy and Other Related
     Properties)
     Section cross-reference(s): 52, 66, 72, 76
     A review with 152 refs. Transient absorption spectroscopy is convenient
AΒ
     to probe the photophys. and photochem. effects of semiconductor
     nanoclusters and heterogeneous charge transfer events at the semiconductor
     interface. The fast kinetic spectroscopy not only enables the detection
     of photogenerated transients in the femtosecond to microsecond time
     domain, but also enables understanding of the photophys. and photochem.
     properties that are unique to semiconductor nanoclusters. For example,
     this technique provided some useful kinetic and mechanistic information of
                                  ***nonlinear***
                                                    optical effects, electro
     size quantization effects,
           ***photochromic***
                                effects, photocurrent
                                                        ***generation***
                         ***charge*** rectification properties of
     photocatalysis and
     semiconductor nanoclusters. These properties are controlled by charge
     sepn., charge trapping and heterogeneous charge transfer at the
     semiconductor interface under UV and visible light irradn. Some recent
     developments in this area are presented.
ST
     transient absorption spectroscopy semiconductor nanocluster review;
     optical transient absorption semiconductor nanocluster review; laser
     transient absorption semiconductor nanocluster review
IT
     Absorption spectra
        (UV and visible; transient absorption spectroscopy of semiconductor
```

```
nanoclusters)
     UV and visible spectra
IT
        (absorption; transient absorption spectroscopy of semiconductor
        nanoclusters)
IT
     Electron transfer
        (from excited dye mols. to semiconductor nanoclusters)
IT
     Clusters
        (nano-; transient absorption spectroscopy of semiconductor
        nanoclusters)
IT
     Catalysts
        (photochem.; transient absorption spectroscopy of semiconductor
        nanoclusters)
IT
     Nanoparticles
         ***Nonlinear*** optical properties
     Quantum size effect
     Semiconductor materials
        (transient absorption spectroscopy of semiconductor nanoclusters)
IT
     Optical absorption
        (transient; transient absorption spectroscopy of semiconductor
        nanoclusters)
    ANSWER 18 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
L5
ΑN
     1996:703842 CAPLUS
DN
     126:8885
     Entered STN: 27 Nov 1996
ED
     Multiple-grating formation in photorefractive polymers with azo-dye
ΤI
     chromophores.
     Smith, M. A.; King, N. R.; Mitchell, G. R.; O'Leary, S. V.
ΑU
     J. J. Thomson Physical Laboratory, University Reading, Reading, RG6 6AF,
CS
     UK
SO
     Proceedings of SPIE-The International Society for Optical Engineering
     (1996), 2850 (Organic Photorefractive Materials and Xerographic
     Photoreceptors), 14-21
     CODEN: PSISDG; ISSN: 0277-786X
     SPIE-The International Society for Optical Engineering
PB
DT
     Journal
     English
LΑ
     36-5 (Physical Properties of Synthetic High Polymers)
CC
    Section cross-reference(s): 73
AΒ
     Photorefractive polymers which incorporate azo-dyes as the
                                                                  ***non***
       ***linear***
                     chromophore element, can be used not only for generating
     gratings by the photorefractive effect, but also by
       ***photoisomerization***
                                 of the azo-dye. In the latter mechanism,
     repeated trans-cis
                          ***isomerization*** causes the chromophore mols. to
     become aligned at right angles to the laser polarization direction,
     thereby making the material birefringent. These two phenomena are to a
     large degree independent, and can be studied sep., by appropriate choice
     of polarization direction of the interacting beams. Furthermore, the
     diffraction efficiency of the photorefractive gratings is a very sensitive
     function of the poling field strength, while that of the
       ***photoisomerization***
                                  gratings is less so. In this work, we
     investigate the components diffracted from each of these grating formed in
     a hybrid photorefractive polymer material PVK:TNF:DEACST:disperse red 1.
     We then explore the possibility of performing some simple optical
     processing applications, exploiting the flexibility provided by this
     multiple grating process. A scheme for producing a novelty filter, which
     displays only the moving parts of a scene is considered.
                                                               The limitations
     of these films for such processing applications are discussed.
ST
     grating multiple photorefractive polymer; azo dye chromophore polymer
     grating
IT
     Diffraction gratings
        (multiple-grating formation in photorefractive polymers with azo-dye
        chromophores)
IT
     Photorefractive materials
     RL: PRP (Properties)
        (multiple-grating formation in photorefractive polymers with azo-dye
       chromophores)
IT
       ***Isomerization***
           ***photoisomerization*** ; multiple-grating formation in
       photorefractive polymers with azo-dye chromophores)
IT
     2872-52-8, Disperse REd1
                                25067-59-8, Poly(N-
                                                    ***vinylcarbazole*** )
     119516-33-5, 4-N, N-Diethylamino-(E)-cinnamonitrile
```

```
RL: PRP (Properties)
   (multiple-grating formation in photorefractive polymers with azo-dye
   chromophores)
ANSWER 19 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
1996:422405 CAPLUS
125:71414
Entered STN: 18 Jul 1996
Photochemistry and Photophysics of Liquid Interfaces by Second Harmonic
Spectroscopy
Eisenthal, K. B.
Department of Chemistry, Columbia University, New York, NY, 10027, USA
Journal of Physical Chemistry (1996), 100(31), 12997-13006
CODEN: JPCHAX; ISSN: 0022-3654
American Chemical Society
Journal; General Review
English
74-0 (Radiation Chemistry, Photochemistry, and Photographic and Other
Reprographic Processes)
Section cross-reference(s): 66
The study of photochem. and photophys. processes at various liq.
interfaces using second harmonic generation methods is described.
the topics discussed are the dynamics of photoinduced structure changes,
      ***transport*** of
                            ***charge***
                                          across an interface, the
rotational motions of interfacial mols., intermol. energy transfer within
the interface, interfacial photopolymn., and photoprocesses at a
semiconductor/liq. interface. A review with 107 refs.
review photochemistry photophysics liq interface; second harmonic
spectroscopy liq interface review
Energy transfer
Interface
Photochemistry
   (photochem. and photophysics of liq. interfaces by second harmonic
   spectroscopy)
Electron exchange and Charge transfer
    ***Isomerization***
Polymerization
   (photochem., photochem. and photophysics of liq. interfaces by second
   harmonic spectroscopy)
          ***nonlinear***
                           property
   (second-harmonic generation, photochem. and photophysics of liq.
   interfaces by second harmonic spectroscopy)
ANSWER 20 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
1995:859398 CAPLUS
123:269408
Entered STN: 17 Oct 1995
  ***Nonlinear***
                  optical properties of organic MIS structures
Charra, Fabrice
DEIN-SPE, Centre d'Etudes de Saclay, Gif-sur-Yvette, F-91 191, Fr.
MCLC S&T, Section B: Nonlinear Optics (1995), 10(1-4), 287-94
CODEN: MCLOEB; ISSN: 1058-7268
Gordon & Breach
Journal; General Review
English
73-0 (Optical, Electron, and Mass Spectroscopy and Other Related
Properties)
An important particularity of org. semiconductors is their strong coupling
between ***charge*** - ***transport*** and optical properties.
applying a voltage on a Metal-Insulator-Semiconductor (MIS) structure
charges can be injected in the semiconductor under well-controlled
conditions. Based on these principles, new mechanisms of electrooptic
coupling and all-optical coupling (i.e.,
                                          ***nonlinear*** optical)
phenomena appear in org. MIS devices. The use of these properties for the
                     ***charge*** - ***transport***
characterization of
                                                        and for
light-modulating devices are discussed. The example of the assocn. in a
MIS structure of electrooptic coupling and photo-cond. is discussed. It
leads to a new type of elec.-controlled
                                         ***photochromic***
                                                               device. A
review with 16 refs.
  ***nonlinear***
                   optical property org MIS review
Electrooptical effect
Optical
         ***nonlinear***
                           property
```

L5 AN

DN

ED

TI

ΑU

CS

so

PB

DT

LA

CC

AB

IT

IT

L5 AN

DN

ED

TI

ΑU

CS SO

PB

DT

LA

CC

AB

ST

TT

```
optical properties and electrooptical effect of
          ***nonlinear***
        org. MIS structures)
ΙŤ
     Optical instruments
                       ***nonlinear***
                                         optical properties and electrooptical
        (modulators,
        effect of org. MIS structures used in modulators)
     ANSWER 21 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
     1995:228102 CAPLUS
     122:200855
DN
ED
     Entered STN: 06 Dec 1994
                            ***generation***
ΤI
     Novel second harmonic
                                                from intermolecular
       ***charge*** -transfer complexes of styrylpyridinium salts in the
     crystalline state
ΑU
     Sakaquchi, Hiroshi; Nagamura, Toshihiko; Kodama, Masahiro; Nishiyama,
     Nobuaki; Sakashita, Hirofumi; Matsuo, Taku
CS
     Crystalline Films Lab., Shizuoka Univ., Hamamatsu, 432, Japan
SO
     Molecular Crystals and Liquid Crystals Science and Technology, Section A:
     Molecular Crystals and Liquid Crystals (1994), 255, 121-9
     CODEN: MCLCE9; ISSN: 1058-725X
PB
     Gordon & Breach
     Journal
DT
LA
     English
     74-1 (Radiation Chemistry, Photochemistry, and Photographic and Other
     Reprographic Processes)
     Section cross-reference(s): 73
     Novel second harmonic generation (SHG) was obsd. from the cryst. powder of
AB
     the ion-pair charge-transfer (CT) complexes consisting of
     1-methyl-4-(4-nitrostyryl)pyridinium as an acceptor and tetraphenylborate
     as a donor. The second harmonic light intensity dramatically decreased
     upon excitation of a CT band due to the photoinduced electron transfer
     from tetraphenylborate to styrylpyridinium moiety. These results strongly
     suggested that the CT transition in the ion-pair complexes would cause
     SHG. The cryst. structure of an ion-pair CT complex of
     1-methyl-4-(4-cyanostyryl)pyridinium tetraphenylborate which also
     exhibited strong SHG was investigated to elucidate the mechanism of the
     SHG in the ion-pair CT complex system.
st
     styrylpyridinium salt charge transfer complex photophys; second harmonic
     generation styrylpyridinium ion pair; ***nonlinear***
                                                             optical property
     styrylpyridinium salt; photochem control second harmonic generation
IT
       ***Photochromism***
        (of intermol. charge-transfer complexes of styrylpyridinium salts in
        cryst. state)
IT
     Crystal structure
        (of ion-pair charge-transfer complex of methyl(cyanostyryl)pyridinium
        tetraphenylborate)
IT
     Electron exchange and Charge transfer
        (photochem., novel second harmonic
                                            ***generation***
                                                              from intermol.
          ***charge*** -transfer complexes of styrylpyridinium salts in cryst.
        state)
TT
              ***nonlinear***
                                property
    Optical
        (second-harmonic generation, novel second harmonic ***generation***
                       ***charge*** -transfer complexes of styrylpyridinium
        from intermol.
        salts in cryst. state)
     161859-28-5, 1-Methyl-4-(4-cyanostyryl)pyridinium tetraphenylborate
IT
     RL: PEP (Physical, engineering or chemical process); PRP (Properties);
    PROC (Process)
        (novel second harmonic generation and crystal structure of)
     140651-15-6, 1-Methyl-4-(4-nitrostyryl)pyridinium tetraphenylborate
IT
     RL: PEP (Physical, engineering or chemical process); PRP (Properties);
    PROC (Process)
        (novel second harmonic ***generation***
                                                    from intermol.
          ***charge*** -transfer complexes of styrylpyridinium salts in cryst.
        state)
IT
     140651-16-7, 1-Methyl-4-(4-nitrostyryl)pyridinium methylbenzenesulfonate
     140651-17-8, 1-Methyl-4-(4-nitrostyryl)pyridinium perchlorate
     RL: PEP (Physical, engineering or chemical process); PRP (Properties);
    PROC (Process)
                               ***generation*** from intermol.
        (novel second harmonic
          ***charge*** -transfer complexes of styrylpyridinium salts in cryst.
        state in relation to)
```

```
DN
    120:165686
     Entered STN: 02 Apr 1994
ED
    Electric field dependence of the laser induced holographic grating
TI
     relaxation of photorefractive polymer
    Wang, C. H.; Ma, R. J.; Zhang, X. Q.; Ducharme, S.; Takacs, J. M.
AU
     Univ. Nebraska, Lincoln, NE, USA
CS
     Report (1992), Order No. AD-A252641, 10 pp. Avail.: NTIS
SO
     From: Gov. Rep. Announce. Index (U. S.) 1992, 92(20), Abstr. No. 256,369
DT
     Report
LA
     English
    37-5 (Plastics Manufacture and Processing)
CC
     Laser-induced holog. grating relaxation (HGR) electrooptical (EO) and
AB
     second harmonic generation (SHG) studies were carried out in a guest/host
     system subjected to an external elec. field with and without the presence
                         ***transport***
            ***hole***
                                            agent. The guest was NNI, which
     of a
                                  ***nonlinear***
                                                    optical coeff., the host
     has a moderate second-order
                       ***hole***
                                       ***transport***
                                                         agent was DEH. In
     was PMMA, and the
                                                  ***isomer***
                                                                  of NNI
     the presence of DEH, the lifetime of the cis
     detected by using the HGR technique is lengthened by the application of an
     elec. field of moderate strength. The field dependence effect is not
     obsd. when DEH is absent in the sample. The result is interpreted as due
                                      ***isomer***
     to the stabilization of the cis
                                                     by the elec. field
     polarized DEH. The data obtained from SHG and EO measurements appear to
     corroborate the HGR result.
     elec field holog grating relaxation; photorefractive PMMA grating
     relaxation laser
     Holography
        (diffraction gratings, photorefractive PMMA, laser induced relaxation
        of, elec. field effect on)
     Diffraction gratings
        (holog., photorefractive PMMA, laser induced relaxation of, elec. field
        effect on)
IT
     Optical materials
           ***nonlinear*** , composite with PMMA, holog. gratings, laser
        induced relaxation of, elec. field effect on)
IT
     9011-14-7, PMMA
     RL: PRP (Properties)
                 ***nonlinear***
                                  optical compd. and
                                                         ***hole***
        (contq.
          ***transport*** agent, elec. field effect on laser-induced holog.
        grating relaxation of)
     ANSWER 23 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
L5
    1993:243976 CAPLUS
AN
     118:243976
DN
ED
     Entered STN: 12 Jun 1993
     The electric field dependence of the laser induced holographic grating
TΙ
     relaxation of a photorefractive polymer
     Wang, C. H.; Ma, R. J.; Zhang, X. Q.; Ducharme, Stephen; Takacs, J. M.
IIA
     Dep. Chem., Univ. Nebraska, Lincoln, NE, 68588-0304, USA
CS
     Proceedings of SPIE-The International Society for Optical Engineering
SO
     (1993), 1775 (Nonlinear Optical Properties of Organic Materials V), 262-70
     CODEN: PSISDG; ISSN: 0277-786X
DT
     Journal
     English
LA
     73-10 (Optical, Electron, and Mass Spectroscopy and Other Related
CC
     Properties)
     Laser induced holog. grating relaxation (HGR), electro-optical (EO) and
AB
     second harmonic generation (SHG) studies were carried out in a guest/host
     system subject to an external elec. field with and without the presence of
        ***hole***
                       ***transport***
                                        agent. The guest is
     1-(4-nitrophenylazo) 2-naphthyl isobutyrate (NNI), which has moderate
     second order ***nonlinear*** optical ( ***NLO*** ) coeff., the host
     is PMMA and the ***hole***
                                    ***transport*** agent is
     p-diethylaminobenzaldehyde diphenylhydrazone (DEH). In the presence of
     DEH, the lifetime of the cis ***isomer*** of NNI detected by using the
     HGR technique is lengthened by the application of an elec. field of
     moderate strength. The field dependence effect is not obsd. when DEH is
     absent in the sample. The result is interpreted as due to the
     stabilization of the cis- ***isomer*** by the elec. field polarized
     DEH. The data obtained from SHG and EO measurements appear to corroborate
```

1994:165686 CAPLUS

the HGR result.

AΝ

```
laser holog grating relaxation photorefractive polymer; elec field
ST
     dependence holog grating relaxation
IT
     Holography
        (grating relaxation of photorefractive polymer, elec. field dependence
        of laser-induced)
IT
     Polymers, properties
     RL: PRP (Properties)
        (holog. grating relaxation of photorefractive, elec. field dependence
        of laser-induced)
TT
     Electrooptical effect
     Optical
              ***nonlinear***
                                 property
        (of photorefractive polymer, elec. field dependence of laser-induced
        holog. grating relaxation in relation to)
IT
     Laser radiation
        (second harmonic generation of, of photorefractive polymer, elec. field
        dependence of laser-induced holog. grating relaxation in relation to)
IT
     Holography
        (diffraction gratings, relaxation of photorefractive polymer, elec.
        field dependence of laser-induced)
     Diffraction gratings
IT
        (holog., relaxation of photorefractive polymer, elec. field dependence
        of laser-induced)
IT
     9011-14-7, Poly(methyl methacrylate)
     RL: USES (Uses)
        (holog. grating relaxation of doped photorefractive, elec. field
        dependence of laser-induced)
IT
     147598-52-5
     RL: USES (Uses)
        (holog. grating relaxation of photorefractive polymer doped with, elec.
        field dependence of laser-induced)
IT
     68189-23-1, p-Diethylaminobenzaldehyde diphenylhydrazone
     RL: USES (Uses)
        (holog, grating relaxation of photorefractive polymer in presence of,
        elec. field dependence of laser-induced)
     ANSWER 24 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
1.5
     1992:215488 CAPLUS
AN
DN
     116:215488
     Entered STN: 31 May 1992
ED
     Photorefractivity in doped
TI
                                  ***nonlinear***
                                                    organic polymers
     Moerner, W. E.; Walsh, C.; Scott, J. C.; Ducharme, S.; Burland, D. M.;
ΑU
     Bjorklund, G. C.; Twieg, R. J.
CS
     Almaden Res. Cent., IBM, San Jose, CA, 95120-6099, USA
     Proceedings of SPIE-The International Society for Optical Engineering
SO
     (1991), 1560 (Nonlinear Opt. Prop. Org. Mater. 4), 278-89
     CODEN: PSISDG; ISSN: 0277-786X
DT
     Journal
LA
     English
     37-5 (Plastics Manufacture and Processing)
CC
     Section cross-reference(s): 38, 73
AB
     The properties of a new class of materials exhibiting the photorefractive
                            ***nonlinear*** org. polymers, are described.
     effect, doped optical
     Photorefraction (at 647.1 nm) is established by a combination of hologram
     erasability, correlation with photocond. and electrooptic response, and
     enhancement by external fields in numerous samples (178-533 .mu.m thick)
            ***nonlinear***
                             epoxy materials doped with
                                                          ***hole***
       ***transport***
                       agents based on p-(diethylamino)benzaldehyde
     diphenylhydrazone. Diffraction efficiencies .ltoreq.0.1% are obsd. at
     bias fields near 100 kV/cm: A useful property of these materials is that
                     ***nonlinear***
     poling of the
                                      chromophores is partially reversible,
     permitting partial control of the grating readout independent of the
     space-charge field formed. The polarization anisotropy of grating readout
     is consistent with the photorefractive mechanism. Two-beam coupling
     measurements of both absorption and index gratings show (a) the absorption
     gratings are .simeq.10 times smaller than the index gratings, and (b) the
     phase shift of the index grating is near 90.degree., which cannot occur
           ***photochromism*** , heating, or any other process except
     photorefractivity.
                               ***nonlinear*** epoxy resin;
ST
     photorefractivity doped
                                                                ***hole***
       ***transport***
                       agent epoxy photorefractivity
IT
     Electrooptical effect
        (of epoxy resins contg.
                                  ***hole***
                                                 ***transport***
                                                                    agents,
```

```
photorefractivity in relation to)
ΙT
     Epoxy resins, properties
     RL: PRP (Properties)
        (photorefractivity of
                                ***nonlinear*** optical, doped with
                         ***transport***
          ***hole***
                                          agents)
IT
     Optical materials
        ( ***nonlinear*** , epoxy resins doped with ***hole***
          ***transport*** agents, photorefractivity of)
                 83890-47-5
IT
     68189-23-1
     RL: PRP (Properties)
        ( ***hole***
                          ***transport***
                                            agents,
                                                     ***nonlinear***
                                                                        optical
        epoxy resins doped with, photorefractivity of)
TT
     125061-60-1
                  128611-17-6
     RL: PRP (Properties)
        (photorefractivity of
                                ***nonlinear***
                                                  optical, doped with
                         ***transport*** agents)
          ***hole***
     ANSWER 25 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
L5
     1992:193573 CAPLUS
AN
DN
     116:193573
ED
     Entered STN: 16 May 1992
     Novel second harmonic
                            ***generation*** from intermolecular
TI
       ***charge*** -transfer complexes of styrylpyridinium tetraphenylborate
     Sakaguchi, Hiroshi; Nagamura, Toshihiko; Matsuo, Taku
ΑU
CS
     Res. Inst. Electron., Shizuoka Univ., Hamamatsu, 432, Japan
     Journal of the Chemical Society, Chemical Communications (1992), (3),
SO
     CODEN: JCCCAT; ISSN: 0022-4936
DT
     Journal
LA
     English
CC
     22-9 (Physical Organic Chemistry)
AΒ
     Strong second harmonic light was obsd. for the first time from intermol.
     ion-pair charge-transfer complexes of nitrostyrylpyridinium
     tetraphenylborate, the intensity of which decreased on irradn. at
     wavelengths longer than 370 nm to induce charge sepn.
     second harmonic generation styrylpyridinium; charge transfer complex
     second harmonic generation
IT
     Electron exchange and Charge transfer
        (in salts of styrylpyridinium, second harmonic generation in relation
        to)
       ***Photochromism***
IT
        (of methyl(nitrostyryl)pyridinium tetraphenylborate)
               ***nonlinear*** property
        (harmonic
                   ***generation***
                                     , second, from
                                                       ***charge*** -transfer
        complexes of styrylpyridinium)
IT
     Electron exchange and Charge transfer
        (photochem., in methyl(nitrostyryl)pyridinium tetraphenylborate)
IT
     140651-15-6
                  140651-16-7 140651-17-8
     RL: PRP (Properties)
        (ion-pair charge-transfer complex, second harmonic generation from)
     ANSWER 26 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
L5
ИA
     1990:580415 CAPLUS
DN
     113:180415
ED
     Entered STN: 09 Nov 1990
ΤI
     Non-equilibrium charge transfer and low-temperature photorefractive
     effects
ΑU
     Nolte, David D.; Glass, Alastair M.
CS
     AT and T Bell Lab., Murray Hill, NJ, 07974, USA
SO
     Optical and Quantum Electronics (1990), 22(Spec. Issue), S47-S60
     CODEN: OQELDI; ISSN: 0306-8919
DT
     Journal
LA
     English
CC
     73-2 (Optical, Electron, and Mass Spectroscopy and Other Related
     Properties)
AΒ
     Defects and
                  ***charge***
                                / ***transport***
                                                    play crit. roles in the
     photorefractive behavior of electrooptic materials. Thus the
     photorefractive effect can be used as a noninvasive optical probe at the
     electronic properties of these materials. Low temp. photorefractive
     spectroscopy has revealed new insight into carrier trapping and deep level
     defects in semiinsulating InP and GaAs.
     nonequil charge transfer gallium arsenide; photorefractive effect indium
```

```
phosphide
   Optical diffraction
TT
        (for iron-doped indium phosphide)
IT
    · Crystal defects
        (in electrooptical materials, photorefractive study of)
IT
     Electron exchange
        (nonequil., in semiinsulating gallium arsenide and indium phosphide)
       ***Photochromism***
IT
        (of chromium-doped gallium arsenide)
ΙT
     Trapping and Traps
        (of current carriers in semiinsulating gallium arsenide and indium
        phosphide)
     Optical materials
IT
        (electro-, photorefractive behavior in)
IT
               ***nonlinear***
     Optical
                                property
        (four-wave mixing, of chromium-doped gallium arsenide)
     Electric current carriers
IT
        (nonequil., in electrooptical materials)
IT
     1303-00-0, Gallium arsenide, properties
     RL: PRP (Properties)
        (nonequil. charge transfer and low temp. photorefractive effects in
        chromium-contq.)
     7439-89-6, Iron, properties
IT
     RL: PRP (Properties)
        (nonequil. charge transfer and low temp. photorefractive effects in
        indium phosphide contg.)
     22398-80-7, Indium phosphide, properties
IT
     RL: PRP (Properties)
        (nonequil. charge transfer in low temp. photorefractive effects in
        iron-contg.)
     7440-47-3, Chromium, properties
     RL: PRP (Properties)
           ***nonlinear***
                             charge transfer and low temp. photorefractive
        effects in gallium arsenide contg.)
     ANSWER 27 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN
1.5
     1967:453488 CAPLUS
AN
     67:53488
DN
     Entered STN: 12 May 1984
ED
     Configuration of vinyl radicals. The generation and trapping of each
                                     ***isomeric*** pair of vinyl radicals
     member of a configurationally
ΑU
     Sargent, G. Dann; Browne, M. Warren
     Amherst Coll., Amherst, MA, USA
CS
     Journal of the American Chemical Society (1967), 89(11), 2788-90
so
     CODEN: JACSAT; ISSN: 0002-7863
DT
     Journal
LA
     English
     22 (Physical Organic Chemistry)
CC
       ***Isomeric*** vinyl free radicals were generated by treating Na
AB
     naphthalenide with cis- and trans-3-chloro-3-hexene at 0.degree. and
     27.degree. using tetrahydrofuran and 1,2-dimethylethane as solvents.
     only detectable products were cis- and trans-3-hexene in 98% yield with
                               predominating. Since these results excluded
                 ***isomer***
     the trans
     the possibility of a single linear sp radical or 2
                                                           ***isomeric***
                               ***nonlinear*** sp2 vinyl free radicals, it tial ***charge*** -transfer reaction
     configurationally stable,
     was concluded that the initial
                            ***nonlinear*** vinyl radical which is capable of
       ***generates*** a
     facile inversion at the radical site, but the radical is trapped before
     complete equilibration with its configurational ***isomer***
     achieved. A mechanism by which the interconverting free radical can be
     trapped is postulated.
     CONFIGURATION VINYL RADICALS; RADICALS VINYL CONFIGURATION; VINYL RADICALS
     CONFIGURATION
IT
     Stereochemistry
        (of naphthylsodium reaction with (E) - and (Z) -3-chloro-3-hexenes)
     3481-12-7
     RL: PRP (Properties)
        (reaction with 3-chloro-3-hexenes, stereochemistry of)
     17226-34-5
                  17226-35-6
     RL: PRP (Properties)
        (reaction with naphthylsodium, stereochemistry of)
     17326-77-1
                 17440-31-2
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RL: PRP (Properties)
(stereochemistry of)

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(FILE 'HOME' ENTERED AT 08:49:00 ON 14 NOV 2005)

FILE 'CAPLUS' ENTERED AT 08:49:10 ON 14 NOV 2005

L1 197907 S (NLO OR NONLINEAR OR (NON(3A)LINEAR) OR HYPERPOLARIZ?)

L2 57950 S (((CHARGE OR HOLE)(3A)(TRANSPORT? OR GENERAT?)) OR CARBAZOLE

L3 1293 S L1 AND L2

369617 S (PHOTOCHROM? OR DIARYLETHANE OR DIHETEROARYLETHANE OR SPIROPY

27 S L3 AND L4

=> log y

L4 L5

COST IN U.S. DOLLARS SINCE FILE TOTAL

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FULL ESTIMATED COST 161.82 162.03

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SINCE FILE TOTAL ENTRY SESSION

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      6 AUG 30 CASREACT - Enhanced with displayable reaction conditions
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      7 SEP 09 ACD predicted properties enhanced in REGISTRY/ZREGISTRY
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 NEWS 9 OCT 04
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                  to core patent offices
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 NEWS 10
                 STN AnaVist workshops to be held in North America
         OCT 13
 NEWS 11
                 New CAS Information Use Policies Effective October 17, 2005
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                 STN(R) AnaVist(TM), Version 1.01, allows the export/download
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                 DIOGENES content streamlined
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         OCT 27
                 EPFULL enhanced with additional content
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              MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
              AND CURRENT DISCOVER FILE IS DATED 13 JUNE 2005
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=> file caplus
COST IN U.S. DOLLARS
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FULL ESTIMATED COST
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we obsd. two-photon photo-orientation of DE in dichroic absorbance. Disperse red one (DR1) was orientationally bleached in PMMA by polarized multi-photon excitation. Anisotropic refractive index changes result from

the \*\*\*nonlinear\*\*\* photobleaching of DR1 was detected by a polarization confocal microscope, and bit-oriented data storage was demonstrated.

- L2 ANSWER 2 OF 14 CAPLUS COPYRIGHT 2005 ACS on STN
- AN 2005:1076285 CAPLUS
- ED Entered STN: 07 Oct 2005
- TI Photoreactive third-harmonic generation via either one- or two-photon excitation in \*\*\*diarylethene\*\*\* -polymethylmethacrylate polymer thin films: theory and experiment
- AU Lin, Jian Hung; Lai, Ngoc Diep; Wang, Wei Lin; Hsu, Chia Chen
- CS National Chung Cheng Univ., Taiwan
- SO Proceedings of SPIE-The International Society for Optical Engineering (2005), 5935(Linear and Nonlinear Optics of Organic Materials V), 262-272 CODEN: PSISDG; ISSN: 0277-786X
- PB SPIE-The International Society for Optical Engineering
- DT Journal
- LA English

AB

- CC 73 (Optical, Electron, and Mass Spectroscopy and Other Related Properties)
  - Photoreactive third-harmonic (TH) generation at 355 nm in

\*\*\*diarylethene\*\*\* - polymethylmethacrylate (DE-PMMA) polymer thin films is obtained by either one- or two-photon excitation. TH intensity generated from a DE-PMMA polymer thin film decreases, when it is pumped by either 325 nm or 442 nm, which change mol. structure of DE mols. from open-form (A form) to closed-form (B form). TH intensity recovers to its original intensity level, after all B form DE mols. return to A form induced by 532 nm or 1064 nm laser irradn. The exptl. results reveal that \*\*\*hyperpolarizability\*\*\* the second-order (.gamma.) of A form mols. may be larger than that of B form mols. Moreover, TH output efficiency is independent of the angle between the pump and probe polarization directions. Those exptl. results were explained by using a photoinduced isomerization theory based angular hole burning and angular redistribution mechanisms for two-dimensional structure of DE mols. including the two-photon absorption effect of B form. The simulation results are consistent with those of optical pumping TH expt.

- L2 ANSWER 3 OF 14 CAPLUS COPYRIGHT 2005 ACS on STN
- AN 2005:297329 CAPLUS
- ED Entered STN: 07 Apr 2005
- TI Optical manipulation of third-harmonic generation via either one- or two-photon excitation in \*\*\*diarylethene\*\*\* -polymethylmethacrylate polymer thin films
- AU Lai, N. D.; Wang, W. L.; Lin, J. H.; Hsu, C. C.
- CS Department of Physics, National Chung Cheng University, Ming Hsiung, Taichung, Taiwan
- SO Applied Physics B: Lasers and Optics (2005), 80(4-5), 569-572 CODEN: APBOEM; ISSN: 0946-2171
- PB Springer GmbH
- DT Journal
- LA English
- CC 73 (Optical, Electron, and Mass Spectroscopy and Other Related Properties)
  AB Optical manipulation of third harmonic (TH) generation in

\*\*\*diarylethene\*\*\* -polymethylmethacrylate (DE-PMMA) polymer thin films is obtained by either one- or two-photon excitation. TH intensity generated from a DE-PMMA polymer thin film decreases, when it is pumped by either 325 nm or 442 nm laser irradn., which changes the mol. structure of DE mols. from an open-form (A form) to a closed-form (B form). TH intensity recovers to its original intensity level when all B form DE mols. return to an A form after being induced by either 532 nm or 1064 nm laser irradn. The exptl. results reveal that the second-order

\*\*\*hyperpolarizability\*\*\* (.gamma.) of the A form mols. may be larger than that of the B form mols. Moreover, TH output efficiency is independent of the angle between the pump and probe polarization directions. This result is attributed to the two-dimensional structure of DE mols.

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- ANSWER 4 OF 14 CAPLUS COPYRIGHT 2005 ACS on STN L2
- 2005:122207 CAPLUS AN
- Entered STN: 11 Feb 2005 ED
- \*\*\*diarylethene\*\*\* for 3D optical ΤI Investigation of novel two-photon data storage
- Zhang, Fushi; Xu, Fengying; Sun, Fan; Zhao, Funqun; Liu, Xuedong; Huang, ΑIJ Wenhao
- Key lab of Organic Optoelectronics & Molecular Engineering of Ministry of CS Education, Department of Chemistry, Tsinghua Univ., Beijing, 100084, Peop. Rep. China
- Proceedings of SPIE-The International Society for Optical Engineering SO (2005), 5643 (Advances in Optical Data Storage Technology), 1-4 CODEN: PSISDG; ISSN: 0277-786X
- SPIE-The International Society for Optical Engineering PB
- DT Journal
- English LA
- 74 (Radiation Chemistry, Photochemistry, and Photographic and Other CC Reprographic Processes)
- 3D optical data storage can benefit from the \*\*\*non\*\*\* - \*\*\*linear\*\*\* AB process of two-photon absorption (TPA). If some chromphores with large TPA cross section are introduced into the \*\*\*diarylethene\*\*\* enhance its two-photon absorptivities, it is expected to be good recording media for 3D optical data storage. The authors have synthesized four diarylperfluorocyclopentene (DPFCP) derivates for the purpose of \*\*\*nonlinear\*\*\* developing two-photon 3D optical data storage. The two-photon absorption (TPA) properties of the compds. were examd., and the reaction and excited state relaxation dynamics after two-photon excitation were investigated. The TPA cross sections of 2-position DPFCPs were measured by means of TPIF (two-photon induced fluorescence) method, and the highest is 48.times.10-50cm4.cntdot.s.cntdot.photo-1.cntdot.mol.-1. The compds. were successfully applied in two-photon 3D optical data storage based on single beam two-photon femtosecond writing and one-photon fluorescence reading. Ten-layer clear images on DPFCP/PMMA film were obtained, and the data storage d. is high to 2.0.times.1011 bits/cm3.
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- L2ANSWER 5 OF 14 CAPLUS COPYRIGHT 2005 ACS on STN
- 2004:508008 CAPLUS AN
- DN 141:61849
- ED Entered STN: 24 Jun 2004
- Compact light-controlled optical waveguide modulator devices and their operation
- IN Fukutome, Masato
- PA Kyocera Corp., Japan
- SO Jpn. Kokai Tokkyo Koho, 9 pp. CODEN: JKXXAF
- DT Patent
- LA Japanese
- ICM G02F001-313
  - C09K009-02; G02F001-361; G02F001-365; G03C001-73
- 73-11 (Optical, Electron, and Mass Spectroscopy and Other Related Properties)
- FAN.CNT 1

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APPLICATION NO.
    PATENT NO.
                       KIND
                              DATE
                                                                DATE
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                              20040624 JP 2002-343009
    JP'2004177623
                                                               20021126
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PRAI JP 2002-343009
                              20021126
CLASS
             CLASS PATENT FAMILY CLASSIFICATION CODES
PATENT NO.
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 JP 2004177623 ICM
                      G02F001-313
               ICS
                       C09K009-02; G02F001-361; G02F001-365; G03C001-73
JP 2004177623 FTERM 2H123/AA00; 2H123/AA10; 2K002/AB09; 2K002/AB12;
                       2K002/BA01; 2K002/CA05; 2K002/DA06; 2K002/DA08;
                       2K002/GA07; 2K002/HA16; 2K002/HA20
    The devices consist of substrates, optical modulators, and optical
AΒ
    waveguides for light irradn. on the optical modulators.
ST
    optical waveguide modulator light irradn control PMMA;
                                                           ***nonlinear***
    optical material PMMA dye modulator; ***diarylethene*** photochromic
    photoresponse org glass modulator
      ***Nonlinear***
                       optical materials
IT
    Optical modulators
    Optical waveguides
       (compact light-controlled optical waveguide modulator devices and their
       operation)
IT
    Photochromic materials
       ( ***diarylethene*** -based; compact light-controlled optical
       waveguide modulator devices and their operation)
IT
    Organic glasses
    RL: DEV (Device component use); USES (Uses)
       (photoresponse; compact light-controlled optical waveguide modulator
       devices and their operation)
    9011-14-7, Poly(methyl methacrylate)
    RL: DEV (Device component use); USES (Uses)
       (compact light-controlled optical waveguide modulator devices and their
       operation)
    7782-41-4, Fluorine, uses
    RL: DEV (Device component use); MOA (Modifier or additive use); USES
     (Uses)
       (dopant; compact light-controlled optical waveguide modulator devices
       and their operation)
    9011-14-7D, Poly(methyl methacrylate), reaction products with dyes
    RL: DEV (Device component use); USES (Uses)
          ***nonlinear*** optical material; compact light-controlled optical
       waveguide modulator devices and their operation)
    ANSWER 6 OF 14 CAPLUS COPYRIGHT 2005 ACS on STN
L2
AN
    2003:686974 CAPLUS
    139:371499
DN
ED
    Entered STN: 03 Sep 2003
    Reversible switching of molecular ***nonlinear*** optical properties
    of photochromic ***diarylethene*** systems
    Bertarelli, C.; Gallazzi, M. C.; Lucotti, A.; Zerbi, G.
ΑU
CS
    Dipartimento di Chimica, Materiali e Ingegneria Chimica "G. Natta",
    Politecnico di Milano, Milan, 20133, Italy
SO
    Synthetic Metals (2003), 139(3), 933-935
    CODEN: SYMEDZ; ISSN: 0379-6779
PB
    Elsevier Science B.V.
DT
    Journal
LA
    English
    73-10 (Optical, Electron, and Mass Spectroscopy and Other Related
CC
    Properties)
AB
    The vibrational contribution to the 3rd-order mol.
      ***hyperpolarizability*** , .gamma.v, of dithienylethenes in the two
    isomeric forms were measured, revealing that a large change in
      ***nonlinear*** optical ( ***NLO*** ) response accompanies the
    photochromic process, thus opening a window on the possibility of
    application of such systems as switching ***NLO*** materials.
    variation of .gamma.v which accompanies the photochromic process is
    strongly dependent on the mol. design and the electronic effect of the
    substituents. Also, the closed form of ***diarylethene*** bearing
    strong electro-donor substituent in 5,5'-positions on thiophene ring
    exhibits large .gamma.v.
    reversible switching ***nonlinear***
                                           optical property
      ***hyperpolarizability***
                                photochromism ***diarylethene***
```

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IT
                  ***nonlinear*** optical properties
    Third-order
         ***hyperpolarizability*** ; reversible switching of mol.
          ***nonlinear***
                          optical properties of photochromic
          ***diarylethene***
                              systems)
    Molecular vibration
IT
                          optical properties
         ***Nonlinear***
    Photochromism
    Raman spectra
        (reversible switching of mol.
                                        ***nonlinear***
                                                          optical properties of
       photochromic
                     ***diarylethene***
                                          systems)
IT
    Optical switching
        (switching; reversible switching of mol.
                                                   ***nonlinear***
                                                                     optical
       properties of photochromic
                                    ***diarylethene***
                                                          systems)
               ***hyperpolarizability***
IT
    Optical
        (third-order; reversible switching of mol.
                                                     ***nonlinear***
                                                                       optical
       properties of photochromic ***diarylethene***
                                                          systems)
                  182003-69-6
                               620990-06-9
IT
     156686-76-9
    RL: PRP (Properties)
        (reversible switching of mol.
                                        ***nonlinear***
                                                          optical properties of
                      ***diarylethene*** systems)
       photochromic
              THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD
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     ANSWER 7 OF 14 CAPLUS COPYRIGHT 2005 ACS on STN
L2
AN
     2003:72901 CAPLUS
DN
     138:254618
ED
     Entered STN: 30 Jan 2003
     Photoswitching of intramolecular magnetic interaction using photochromic
TI
     compounds
ΑU
     Matsuda, Kenji; Irie, Masahiro
     Graduate School of Engineering, Kyushu University, Higashi-ku, Fukuoka,
CS
     812-8581, Japan
     Springer Series in Chemical Physics (2003), 70 (Chemistry of Nanomolecular
SO
     Systems), 25-40
     CODEN: SSCPDA; ISSN: 0172-6218
PB
     Springer-Verlag
DT
     Journal; General Review
LA
     English
CC
     22-0 (Physical Organic Chemistry)
     Section cross-reference(s): 77
     A review. Recent advances in org. material chem. have helped to realize
AB
     several novel org. functional materials, including org. conductors,
                                    optical materials. One of the current
                   ***nonlinear***
     magnets, and
     frontiers in this field is mol. electronics, where a single mol. plays the
     role of one component in an elec. circuit. For the realization of this
     goal, highly integrated materials with multiple functions are necessary.
     Photochromism is the light-induced reversible transformation of chem.
     species between two isomers that have different absorption spectra.
     two isomers differ from each other not only in the absorption spectra but
     also in various phys. and chem. properties. These property changes can,
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in principle, be utilized to control the function of org. materials. org. radicals placed at the two edges of a .vpi.-conjugative mol. interact magnetically through the framework via the exchange interaction J. Therefore photocontrol of the magnetism of the system is possible when the photochromic moiety is used as a backbone. In this chapter we will describe studies of the photoswitching of the intramol. magnetism by incorporating two radical moieties into a photochromic spin coupler. Photoswitching using a \*\*\*diarylethene\*\*\* \*\*\*diarylethene\*\*\* dimer, which showed similar elec.-circuit behavior, will also be discussed. review photoswitching intramol magnetic interaction photochromic compd Magnetism (intramol.; photoswitching of intramol. magnetic interaction using photochromic compds.) Radicals, properties RL: PRP (Properties) (moieties; photoswitching of intramol. magnetism by incorporating two radicals into a photochromic \*\*\*diarylethene\*\*\* spin coupler) Optical switching (photoswitching of intramol. magnetic interaction using photochromic compds.) Exchange interaction Logic circuits Nanostructures Photochromism (photoswitching of intramol. magnetism by incorporating two radicals into a photochromic \*\*\*diarylethene\*\*\* spin coupler) Nitroxides RL: PRP (Properties) (photoswitching of intramol. magnetism by incorporating two radicals into a photochromic \*\*\*diarylethene\*\*\* spin coupler) RE.CNT THERE ARE 45 CITED REFERENCES AVAILABLE FOR THIS RECORD (1) Anon; Chem Rev 2000, V100(5) (2) Atassi, Y; J Phys Chem 1995, V99, P16320 CAPLUS (3) Balzani, V; Angew Chem, Int Ed Engl 2000, V39, P3349 (4) Blank, M; Science 1981, V214, P70 CAPLUS (5) Bleany, B; Proc R Soc London A 1952, V214, P451 (6) Borden, W; J Am Chem Soc 1977, V99, P4587 CAPLUS (7) Briere, R; Bull Soc Chim France 1965, V11, P3290 (8) Brown, G; Photochromism 1971 (9) Durr, H; Photochromism: Molecules and Systems 1990 (10) Frank, N; J Am Chem Soc 2000, V122, P2053 CAPLUS (11) Gilat, S; J Chem Soc, Chem Commun 1993, P1439 CAPLUS (12) Glarum, S; J Chem Phys 1967, V47, P1374 CAPLUS (13) Glat, S; Chem Eur J 1995, V1, P275 (14) Ichimura, K; Langmuir 1988, V4, P1214 CAPLUS (15) Ikeda, T; Nature 1993, V361, P428 CAPLUS (16) Irie, M; Adv Polym Sci 1990, V94, P27 CAPLUS (17) Irie, M; Bull Chem Soc Jpn 1998, V71, P985 CAPLUS (18) Irie, M; Chem Rev 2000, V100, P1685 CAPLUS (19) Irie, M; J Am Chem Soc 1985, V107, P1024 CAPLUS (20) Irie, M; Macromolecules 1981, V14, P262 CAPLUS (21) Irie Adv, M; Polym Sci 1993, V110, P49 (22). Iwamura, H; Acc Chem Res 1993, V26, P346 CAPLUS (23) Jacobs, S; J Am Chem Soc 1993, V115, P1744 CAPLUS (24) Joachim, C; Nature 2000, V408, P541 CAPLUS (25) Kahn, O; Molecular Magnetism 1993 (26) Kawai, T; Chem Lett 1999, P905 CAPLUS (27) Lehn, J; Supramolecular Chemistry: Concepts and Perspectives 1995 (28) Matsuda, K; Chem Eur J 2001, V7, P3466 CAPLUS (29) Matsuda, K; Chem Lett 2000, P16 CAPLUS (30) Matsuda, K; Chem Lett 2001, P436 CAPLUS (31) Matsuda, K; J Am Chem Soc 1997, V119, P7412 CAPLUS (32) Matsuda, K; J Am Chem Soc 1997, V119, P8058 CAPLUS (33) Matsuda, K; J Am Chem Soc 2000, V122, P7195 CAPLUS (34) Matsuda, K; J Am Chem Soc 2000, V122, P8309 CAPLUS (35) Matsuda, K; J Am Chem Soc 2001, V123, P9896 CAPLUS (36) Matsuda, K; J Org Chem 2001, V66, P8799 CAPLUS (37) Matsuda, K; Tetrahedron Lett 2000, V41, P2577 CAPLUS (38) Matsumoto, T; J Am Chem Soc 1992, V114, P9952 CAPLUS (39) Mitsumori, T; J Am Chem Soc 1995, V117, P2467 CAPLUS

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     ANSWER 8 OF 14 CAPLUS COPYRIGHT 2005 ACS on STN
L2
     2002:255330 CAPLUS
AN
DN
     137:33571
     Entered STN: 05 Apr 2002
ED
     Radical copolymerization behavior of a highly fluorinated cyclic olefin
ΤI
     with vinyl ether
     Nishida, Shozo; Nishida, Haruo; Endo, Takeshi
AU
     Tsukuba Research Laboratories, JSR Corporation, Tsukuba, 305-0841, Japan
CS
     Journal of Polymer Science, Part A: Polymer Chemistry (2002), 40(8),
SO
     CODEN: JPACEC; ISSN: 0887-624X
PB
     John Wiley & Sons, Inc.
DT
     Journal
LA
     English
     35-3 (Chemistry of Synthetic High Polymers)
CC
AΒ
     The copolymn. of a highly fluorinated cyclic monomer,
     octafluorocyclopentene (OFCPE, M1), with Et vinyl ether (EVE, M2) was
     investigated with a radical initiator in bulk. Despite the poor
     homopolymerizability of each monomer, the copolymn. proceeded
     successfully, and the mol. wts. of the copolymers were more than 10,000.
     Incorporation of the OFCPE units into the copolymer led to an increase in
     the glass-transition point. The copolymer compn. was detd. from 1H NMR
     spectra and elemental anal. data. The molar fraction of the OFCPE unit in
     the copolymer increased and approached but did not exceed 0.5.
     monomer reactivity ratios were estd. by the Yamada-Itahashi-Otsu
       ***nonlinear***
                        least-squares procedure as r1,OFCPE = -0.008 .+-. 0.010
     and r2, EVE = 0.192 .+-. 0.015. The reactivity ratios suggest that the
     copolymn. may proceed in an alternating manner in the case of an excess
     feed of OFCPE.
ST
       ***perfluorocyclopentene***
                                     reactivity ratio radical polymn vinyl
     ether
IT
     Fluoropolymers, preparation
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (radical prepn. and glass transition temp. of)
IT
     Reactivity ratio in polymerization
                      ***perfluorocyclopentene***
                                                    with Et vinyl ether)
        (radical; of
IT
     363593-48-0P, Ethyl vinyl ether- ***perfluorocyclopentene***
     RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
        (radical prepn. and glass transition temp. of)
     559-40-0, Octafluorocyclopentene
     RL: PRP (Properties); RCT (Reactant); RACT (Reactant or reagent)
        (reactivity ratio in radical polymn. with Et vinyl ether)
IT
     109-92-2, Ethyl vinyl ether
     RL: PRP (Properties); RCT (Reactant); RACT (Reactant or reagent)
        (reactivity ratio in radical polymn. with ***perfluorocyclopentene***
              THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
       12
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L2
AN
     2001:792858 CAPLUS
DN
     136:77164
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ED

Entered STN: 01 Nov 2001

```
Novel photoresponsive polymer based on
ΑU
     Kawai, Tsuyoshi; Irie, Masahiro
     Dep. Applied Chem., Fac. Eng., Kyushu Univ., Hakozaki 6-10-1, Higashi-ku,
CS
     Fukuoka, 812-8581, Japan
     Denki Zairyo Gijutsu Zasshi (2000), 9(2), 204-207
SO
     CODEN: DZGZFE; ISSN: 0918-9890
     Denki Zairyo Gijutsu Kondankai
PB
DT
     Journal
LA
     English
     74-9 (Radiation Chemistry, Photochemistry, and Photographic and Other
CC
     Reprographic Processes)
     Section cross-reference(s): 36
     Conjugated polymer based on the bis(benzothiophen-3-yl)
AΒ
       ***perfluorocyclopentene***
                                    can exhibit drastic changes of optical and
     elec. properties upon photochromic reactions of the
                                                         ***diarylethene***
     units. A novel photoresponsive polymer based on ***diarylethene***
     and dialkylfluorene was prepd. The polymer was prepd. by Ru-complex
     catalyzed coupling reaction of the two components. The quartz glass
     supported polymer film had strong absorption band at .apprx.334 nm. After
    UV exposure the pale yellow film turned to violet showing new absorption
     band at 560 nm, which is characteristic band for the ring-closed form of
           ***diarylethene***
                              sites. After the irradn. with the visible
     the
     light .lambda. >480 nm the original yellow film was recovered and the
     absorption band at 560 nm disappeared. This photochromic cycle was
     repeated for more than 20 cycles and no marked degrdn. was obsd. The max.
     conversion of the photochromic site was .apprx.40 % in soln. and .apprx.14
     % in solid film. The polymer had photoluminescence max. at .apprx.410 nm
     which decreased rapidly with exposure. After.apprx.5 min., the emission
     was totally quenched, at the same time only 14% of the diarylene sites was
     cyclized even at the photostationary state. This remarkable
                                                                   ***non***
                      phenomenon can be explained by the term of the efficient
        ***linear***
     excitation energy transfer in the polymer chain. That is, the small amt.
     of the ring-closed form sites which have second absorption band at about
     400 nm acts efficiently as the energy acceptor and the emission quencher.
     The cond. of the polymer also changed with exposure. The cond. was about
     1.2x10-12 S-cm-1 in the 35 % cyclized state photostationary state achieved
     in the soln. phase, and it decreased to 5.3 x 10-13 S-cm-1 upon irradn.
     with visible light. The I-V characteristics of the Al/polymer/ITO type
     junction device were studied. In this device the Al electrode was evapd.
     onto the polymer film on the ITO electrode. In the original state, the
     I-V curve showed clear current rectification and the forward current was
     obsd. when pos. bias was applied to the ITO electrode. After the UV light
     irradn. the reverse biased current was found to increase. That is,
     electronic state of the ring-closed form
                                               ***diarylethene***
     inter-gap state in the junction characteristics at Al-polymer interface
     and photochromic reaction in small amt. of the ***diarylethene***
     sites seems to results in the drastic increase in the leaking reverse
     direction current.
     photochromism photocond elec cond
                                        ***diarylethene***
     photopolymer; benzothiophenylperfluorocyclopentene based polymer
     photochromism photocond elec cond
     Polymers, properties
IT
     RL: PEP (Physical, engineering or chemical process); PNU (Preparation,
     unclassified); PRP (Properties); PREP (Preparation); PROC (Process)
        (conjugated; photochromism and luminescence and cond. of
        photoresponsive polymer based on bis(benzothiophenyl)
          ***perfluorocyclopentene***
IT
     Luminescence quenching
     Photoinduced energy transfer
        (intramol.; photochromism and luminescence and cond. of photoresponsive
        polymer based on bis(benzothiophenyl) ***perfluorocyclopentene*** )
IT
     Ring opening
        (photochem.; photochromism and luminescence and cond. of
        photoresponsive polymer based on bis(benzothiophenyl)
          ***perfluorocyclopentene***
IT
     Electric conductivity
     Electric current-potential relationship
     Electrochromic devices
     Luminescence
     Photochromic materials
     Photochromism
     Photoconductivity
```

\*\*\*diarylethene\*\*\*

ΤI

```
(photochromism and luminescence and cond. of photoresponsive polymer
        based on bis(benzothiophenyl) ***perfluorocyclopentene***
IT
     Cyclization
        (photocyclization; photochromism and luminescence and cond. of
       photoresponsive polymer based on bis(benzothiophenyl)
          ***perfluorocyclopentene***
     384340-52-7P
IT
    RL: PEP (Physical, engineering or chemical process); PNU (Preparation,
    unclassified); PRP (Properties); PREP (Preparation); PROC (Process)
        (photochromism and luminescence and cond. of photoresponsive polymer
       based on bis(benzothiophenyl) ***perfluorocyclopentene*** )
              THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT 6
RE
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(5) Koshido, T; J Phys Chem 1995, V16, P6110
(6) Koshido, T; Synth Met 1995, V73, P257 CAPLUS
    ANSWER 10 OF 14 CAPLUS COPYRIGHT 2005 ACS on STN
L2
    2000:764182 CAPLUS
ΑN
DN
    134:92993
ED
    Entered STN: 01 Nov 2000
ΤI
       ***Nonlinear***
                                                 ***diarylethenes***
                       optical properties of
ΑIJ
    Delaire, J. A.; Fanton-Maltey, I.; Chauvin, J.; Nakatani, K.; Irie, M.
    PPSM, UMR 8531 of CNRS, Ecole Normale Superieure de Cachan, Cachan, 94235,
CS
    Fr.
so
    Molecular Crystals and Liquid Crystals Science and Technology, Section A:
    Molecular Crystals and Liquid Crystals (2000), 345, 233-238
    CODEN: MCLCE9; ISSN: 1058-725X
PB
    Gordon & Breach Science Publishers
DT
    Journal
LA
    English
    73-10 (Optical, Electron, and Mass Spectroscopy and Other Related
CC
     Properties)
     Section cross-reference(s): 22
     Products .mu..beta. of the 1st order
                                            ***hyperpolarizabilities***
AB
     .beta. of both isomers of dithienyl ethene derivs. by their ground state
     dipole moments .mu. were detd. exptl. by the EFISHG technique, then calcd.
    by a semi-empirical method. Both calcns. and measurements put into
     evidence a large increase in .mu..beta. in going from the open isomer (OF)
     to the largely .pi. conjugated closed isomer (CF). This photoinduced
     change in 2nd order ***NLO***
                                       coeff. was used to modulate the Second
    Harmonic Generation (SHG) signal generated by a PMMA film doped with one
       ***diarylethene***
                           under alternate irradn. with UV (325 nm) and visible
     (514 nm) light.
                                            ***diarylethene***
ST
       ***nonlinear***
                         optical property
        (dipole moment for;
                             ***nonlinear***
                                                optical properties of
          ***diarylethenes*** )
IT
     Cyclization
        (effect of;
                      ***nonlinear***
                                        optical properties of
          ***diarylethenes*** )
ΙT
    Dipole moment
        (ground state;
                         ***nonlinear***
                                           optical properties of
          ***diarylethenes*** )
IT
       ***Nonlinear***
                       optical materials
         ***Nonlinear***
                          optical properties
              ***hyperpolarizability***
     Second-harmonic generation
          ***nonlinear***
                            optical properties of
                                                     ***diarylethenes*** )
IT
     9011-14-7, PMMA
    RL: NUU (Other use, unclassified); USES (Uses)
                  ***nonlinear***
                                    optical properties of
          ***diarylethenes*** )
IT
     112440-47-8
                 115755-82-3
                                 168688-69-5
                                               185202-32-8
    RL: PRP (Properties)
           ***nonlinear***
                             optical properties of
                                                     ***diarylethenes*** )
        (
RE.CNT
             THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS RECORD
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L2
     1999:676500 CAPLUS
AN
DN
     132:257359
     Entered STN: 25 Oct 1999
ED
ΤI
     Temporal stability of azo second-order
                                              ***nonlinear***
                                                                optical
     chromophores linked with perfluorocyclopentenyl moiety
     Matsui, Masaki; Muramatsu, Hiroshige; Funabiki, Kazumasa; Shibata,
ΑU
     Katsuyoshi; Hirota, Kazuo; Hosoda, Masahiro; Tai, Kazuo
     Department of Chemistry, Faculty of Engineering, Gifu University,
CS
     Yanagido, Gifu, 501-1193, Japan
     Shikizai Kyokaishi (1999), 72(8), 489-493
SO
     CODEN: SKYOAO; ISSN: 0010-180X
PB
     Shikizai Kyokai
DT
     Journal
     English
LA
     73-2 (Optical, Electron, and Mass Spectroscopy and Other Related
CC
     Properties)
     Section cross-reference(s): 36, 37
     The temporal stability of azo second-order
                                                  ***nonlinear***
AΒ
                                                                    optical
     chromophores linked with a perfluorocyclopentenyl moiety doped in polymers
     was examd. The retention of second-order ***nonlinear***
                                                                   optical
     coeff. (d33) of 1,2-bis[4-[4-[N-ethyl-N-(2-hydroxyethyl)amino] phenylazo]
                   ***perfluorocyclopentene***
     phenylthio]
                                                doped in poly(Me methacrylate)
     after 530 h at 50.degree.C was 5 %. The retention of second-order
                        optical coeff. (d33) of 1,2-bis[4-(2,4-
       ***nonlinear***
     diaminophenylazo)phenylthio] ***perfluorocyclopentene***
     poly (Me methacrylate-co-methacrylic acid) was improved to 45wt.% by the
     amino formation between azo dye and polymer matrix.
                             ***nonlinear***
ST
     temporal stability azo
                                               optical chromophore; doped
     polymer azo
                   ***nonlinear***
                                     optical chromophore temporal stability
IT
     Azo dyes
     Chromophores
     Second-order
                    ***nonlinear***
                                      optical properties
     Thermal stability
        (prepn. and temporal stability of neat and polymer-doped azo
        second-order ***nonlinear***
                                         optical chromophores linked with
       perfluorocyclopentenyl moiety)
IT
       ***Nonlinear*** optical materials
        (second-order; prepn. and temporal stability of neat and polymer-doped
        azo second-order
                          ***nonlinear***
                                             optical chromophores linked with
        perfluorocyclopentenyl moiety)
IT
     178401-05-3P
                    262858-27-5P
                                   262858-28-6P
                                                  262858-29-7P
                                                                 262858-30-0P
     RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
        (chromophore; prepn. and temporal stability of neat and polymer-doped
                          ***nonlinear***
                                             optical chromophores linked with
        azo second-order
        perfluorocyclopentenyl moiety)
IT
     9011-14-7, PMMA
                      25086-15-1, Methacrylic acid-methyl methacrylate
     copolymer
     RL: PRP (Properties)
        (prepn. and temporal stability of neat and polymer-doped azo
                      ***nonlinear***
                                        optical chromophores linked with
        second-order
        perfluorocyclopentenyl moiety)
IT
     243461-38-3
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (starting material; prepn. and temporal stability of neat and
        polymer-doped azo second-order
                                         ***nonlinear***
                                                           optical chromophores
        linked with perfluorocyclopentenyl moiety)
              THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
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     ANSWER 12 OF 14 CAPLUS COPYRIGHT 2005 ACS on STN
L2
AN
     1999:405855 CAPLUS
DN
     131:215636
     Entered STN: 01 Jul 1999
ED
TI
     Synthesis of azo chromophores containing a perfluorocyclo-alkenyl moiety
     and their second-order optical nonlinearity
     Matsui, Masaki; Tsuge, Michinori; Funabiki, Kazumasa; Shibata, Katsuyoshi;
AU
     Muramatsu, Hiroshige; Hirota, Kazuo; Hosoda, Masahiro; Tai, Kazuo;
     Shiozaki, Hisayoshi; Kim, Misa; Nakatsu, Kazumi
     Faculty of Engineering, Department of Chemistry, Gifu University, Gifu,
CS
     501-1193, Japan
     Journal of Fluorine Chemistry (1999), 97(1-2), 207-212
SO
     CODEN: JFLCAR; ISSN: 0022-1139
PR
    Elsevier Science S.A.
DT
     Journal
LA
     English
     41-3 (Dyes, Organic Pigments, Fluorescent Brighteners, and Photographic
CC
     Sensitizers)
     Section cross-reference(s): 38, 73
     1,2-Bis[4-[4-[N-ethyl-N-(2-hydroxyethyl)amino]phenylazo]phenylthio]
AΒ
       ***perfluorocyclopentene*** , having two independent intramol. push-pull
     chromophores in the mol., showed a higher second-order ***nonlinear***
     coeff. (d33) than its perfluorocyclobutenyl, perfluorocyclohexenyl, and
     nitroperfluorocyclopentenyl analogs in PMMA or polycarbonate.
               ***nonlinear***
                                 optical prepn
ST
     azo dye
IT
     Bond length
                                          ***nonlinear***
        (carbon-carbon; in second-order
                                                             optical azo dye)
IT
     Bond length
        (carbon-nitrogen; in second-order
                                             ***nonlinear***
                                                               optical azo dye)
IT
     Bond length
                                                             optical azo dye)
        (carbon-sulfur; in second-order
                                           ***nonlinear***
TT
     Bond angle
        (in second-order
                           ***nonlinear***
                                             optical azo dye)
     Crystal structure
        (of second-order
                           ***nonlinear***
                                             optical azo dye)
IT
     Azo dyes
                                  ***nonlinear***
        (prepn. of second-order
                                                     optical)
                    ***nonlinear*** optical properties
IT
        (prepn. of second-order ***nonlinear***
                                                    optical azo dyes)
     Polycarbonates, uses
IT
     RL: NUU (Other use, unclassified); USES (Uses)
        (second-order optical nonlinearity of azo dyes in)
IT
     Bond angle
        (torsional; in second-order
                                      ***nonlinear***
                                                         optical azo dye)
     92-50-2, 2-(N-Ethylanilino)ethanol
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (coupling component; prepn. of second-order
                                                       ***nonlinear***
                                                                         optical
        azo dyes)
     336-19-6, 1,2-Dichloroperfluorocyclohexene
                                                   377-93-5,
     1,2-Dichloroperfluorocyclobutene 706-79-6, 1,2-
     Dichloroperfluorocyclopentene
                                    1193-02-8, 4-Aminothiophenol
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (diazo component starting material; prepn. of second-order
          ***nonlinear***
                            optical azo dyes)
     243461-36-1P, 1,2-Bis (4-aminophenylthio) perfluorocyclobutene
IT
     243461-38-3P, 1,2-Bis(4-aminophenylthio) ***perfluorocyclopentene***
     243461-39-4P, 1,2-Bis (4-aminophenylthio) perfluorocyclohexene
     243461-40-7P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
                                                    ***nonlinear***
                                                                      optical
        (diazo component; prepn. of second-order
        azo dyes)
IT
     178401-04-2P
                    178401-05-3P
                                   178401-06-4P
                                                   243461-41-8P
     RL: PRP (Properties); SPN (Synthetic preparation); TEM (Technical or
     engineered material use); PREP (Preparation); USES (Uses)
        (dye; prepn. of second-order
                                       ***nonlinear***
                                                         optical azo dyes)
     19433-93-3, 5-(Acetylamino)-N, N-diethyl-2-methoxyaniline
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
```

```
(model compd. coupling component; prepn. of second-order
                           optical azo dyes)
          ***nonlinear***
IT
     81924-79-0P, 5-Acetamido-N, N-diethyl-2-methoxy-4-(4-nitrophenylazo) aniline
     243461-29-2P, 5-Acetamido-N, N-diethyl-2-methoxy-4-(phenylazo) aniline
     243461-30-5P, 5-Acetamido-4-(4-chlorophenylazo)-N,N-diethyl-2-
                      243461-31-6P, 5-Acetamido-N, N-diethyl-4-(4-
    methoxyaniline
                                         243461-33-8P, 5-Acetamido-N, N-diethyl-
     fluorophenylazo) -2-methoxyaniline
     2-methoxy-4-[4-(trifluoromethyl)phenylazo]aniline
                                                         243461-34-9P
     243461-35-0P
    RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
        (model compd.; prepn. of second-order
                                                ***nonlinear***
        dyes)
IT
     4485-08-9, 4-Nitronitrosobenzene
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (prepn. of second-order
                                  ***nonlinear***
                                                    optical azo dyes)
TT
     9011-14-7, PMMA
    RL: NUU (Other use, unclassified); USES (Uses)
        (second-order optical nonlinearity of azo dyes in)
              THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
RE
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     ANSWER 13 OF 14 CAPLUS COPYRIGHT 2005 ACS on STN
L2
     1999:54878 CAPLUS
AN
DN
     130:182943
     Entered STN: 27 Jan 1999
ED
     Reversible photoinduced modifications of polymers doped with photochromes
     : anisotropy, photo-assisted poling and surface gratings
     Atassi, Yomen; Chauvin, Jerome; Delaire, Jacques; Delouis, Jean-Francois;
     Fanton-Maltey, Isabelle; Nakatani, Keitaro
     PPSM, URA 1906 of CNRS, Ecole Normale Supdrieure de Cachan, Cachan, 94235,
CS
     Molecular Crystals and Liquid Crystals Science and Technology, Section A:
SO
     Molecular Crystals and Liquid Crystals (1998), 315, 313-324
     CODEN: MCLCE9; ISSN: 1058-725X
PB
     Gordon & Breach Science Publishers
DT
     Journal; General Review
LA
     English
     36-0 (Physical Properties of Synthetic High Polymers)
CC
     Section cross-reference(s): 73, 76
     In this paper, different optical methods which lead to orientation or
AΒ
     translation of several photochromes in polymer matrixes are reviewed with
     17 refs. These methods lead to dichroic or birefringent, or
                         optically active materials or to stable surface
       ***nonlinear***
     gratings. They have mainly been demonstrated for azobenzene-type
     photochromes, but some of these effects also occur for other kinds of
     photochromes like spiropyrans, fulgides or
                                                  ***diarylethenes***
     reversible photoinduced modification photochrome doped polymer review;
ST
     anisotropy photochrome doped polymer reversible modification review;
     poling photochrome doped polymer reversible modification review; surface
     grating photochrome doped polymer reversible modification review
IT
     Molecular orientation
     Optical anisotropy
     Second-harmonic generation
        (anisotropy, photo-assisted poling and surface gratings of reversible
        photoinduced modified PMMA doped with photochromes)
IT
     Polymers, properties
```

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RL: POF (Polymer in formulation); PRP (Properties); TEM (Technical or
    engineered material use); USES (Uses)
       (anisotropy, photo-assisted poling and surface gratings of reversible
        photoinduced modified PMMA doped with photochromes)
     Chromophores
        (photo-; anisotropy, photo-assisted poling and surface gratings of
        reversible photoinduced modified PMMA doped with photochromes)
    Optical switching
        (photoinduced; anisotropy, photo-assisted poling and surface gratings
       of reversible photoinduced modified PMMA doped with photochromes)
     Isomerization
        (photoisomerization; anisotropy, photo-assisted poling and surface
        gratings of reversible photoinduced modified PMMA doped with
        photochromes)
    Diffraction gratings
        (surface; anisotropy, photo-assisted poling and surface gratings of
        reversible photoinduced modified PMMA doped with photochromes)
     9011-14-7, PMMA
     RL: PRP (Properties)
        (anisotropy, photo-assisted poling and surface gratings of reversible
        photoinduced modified PMMA doped with photochromes)
              THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
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(2) Chauvin, J; SPIE 1997, V2998, P205 CAPLUS
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    ANSWER 14 OF 14 CAPLUS COPYRIGHT 2005 ACS on STN
     1995:762770 CAPLUS
    123:339588
    Entered STN: 29 Aug 1995
    Light-triggered molecular devices: photochemical switching of optical and
    electrochemical properties in molecular wire type
                                                         ***diarylethene***
     species
    Gilat, Sylvain L.; Kawai, Stephen H.; Lehn, Jean-Marie
    Chim. Interactions Mol., Coll. France, Paris, 75005, Fr.
    Chemistry--A European Journal (1995), 1(5), 275-84
                                                            Published in:
    Angew. Chem., Int. Ed. Engl., 34(15)
     CODEN: CEUJED; ISSN: 0947-6539
    VCH
    Journal
    English
     27-8 (Heterocyclic Compounds (One Hetero Atom))
/ Structure 1 in file .gra /
    Bispyridine I (R = 4-pyridyl) and the corresponding bispyridinium were
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synthesized as the uncyclized isomers from 3,5-dibromo-2-methylthiophene in overall yields of 43 and 44%, resp. The \*\*\*diarylethene\*\*\* photochromes I [R = 1,3-benzodithiol-2-yl, CHO, CH:C(CN)2, etc.], substituted with electron donors and acceptors, were prepd. from 5-methylthiophene-2-carboxyaldehyde in 21-32% overall yield. All of the compds. were found to exhibit pronounced photochromic properties. Irradn. with UV light resulted in essentially complete photocyclization of the

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be reconverted back to the open state with visible light of .lambda. > 600
         The absorption maxima of the described compds. in their closed forms
    are shifted far towards, and even into, the near-IR region. Whereas no
    thermochromic properties were obsd. for the open isomers, the rates of
    thermal decoloration of the cyclized forms was found to be highly
    dependent on the nature of the substituents on the thiophene rings. It
    was demonstrated that reversible photochem. interconversion between the
    two photochromic states could be used to effectively switch a no. of phys.
    properties. Thus, the bispyridinium formed from I (R = 4-pyridyl) and I
    (R = 1,3-benzodithiol-2-yl) represent two kinds of redox switches, the
    former in redn. and the latter in oxidn., in which electron conduction is
    switched on in the closed state and off in the open state. I (R =
    1,3-benzodithiol-2-yl) may also be considered to be a photoswitchable
    analog of tetrathiafulvalene type substances. On the other hand, I (R =
    2-benzodithiolyl) displays a marked increase in
                                                       ***nonlinear***
    optical activity on conversion from the open to the closed form. Such
    systems are prototypes of photoswitchable mol. wires where electron
    conduction and push-pull interaction can be reversibly modulated by an
    external stimulus, namely, irradn. by light.
    redox switch
                   ***diarylethene***
                                         deriv; photochem switching optical
    electrochem property; elec cond photochem switching;
                                                            ***nonlinear***
    optical activity photochem switching; thermochromism
                                                            ***diarylethene***
    deriv; photochromism ***diarylethene***
                                                 deriv;
    thienylperfluorocyclopentene photochem switching;
       ***perfluorocyclopentene*** dithienyl photochem switching; mol wire
    photoswitchable prototype
    Thermochromism
        (of closed isomers of bis(thienyl)ethene compds.)
    Electric conductivity and conduction
              ***nonlinear***
    Optical
                                property
    Photochromism
        (photochem. switching of optical and electrochem. properties in mol.
       wire type bis(thienyl)ethene species)
                                                  154566-76-4P
                                                                 154566-77-5P
                                   154566-75-3P
    154566-71-9P
                   154566-73-1P
                                                                 154566-82-2P
                                                  154566-81-1P
                                   154566-80-0P
    154566-78-6P
                   154566-79-7P
                                   170698-91-6P
                                                  170698-92-7P
    154566-83-3P
                   154566-84-4P
    RL: PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); PREP
     (Preparation); RACT (Reactant or reagent)
        (photochem. switching of optical and electrochem. properties in mol.
       wire type bis(thienyl)ethene species)
                                           ***Perfluorocyclopentene***
    109-77-3, Malononitrile
                               559-40-0,
                               62217-34-9
                                            74129-11-6, 4-Bromopyridine
    13679-70-4
                 29421-73-6
    hydrobromide
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (photochem. switching of optical and electrochem. properties in mol.
        wire type bis(thienyl)ethene species)
                                                 154566-74-2P
                                                                154586-93-3P
    29421-75-8P
                   154566-69-5P
                                  154566-70-8P
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (photochem. switching of optical and electrochem. properties in mol.
        wire type bis(thienyl)ethene species)
=> s l1 and (((charge or hole)(2w)(transport or generat?)) or polyvinylcarbazole or carbazole or v
       484684 CHARGE
         62432 CHARGES
        519427 CHARGE
                 (CHARGE OR CHARGES)
        188697 HOLE
        110661 HOLES
        253252 HOLE
                 (HOLE OR HOLES)
        671926 TRANSPORT
          5403 TRANSPORTS
        674068 TRANSPORT
                 (TRANSPORT OR TRANSPORTS)
        925147 GENERAT?
         27522 (CHARGE OR HOLE) (2W) (TRANSPORT OR GENERAT?)
          2223 POLYVINYLCARBAZOLE
            34 POLYVINYLCARBAZOLES
          2250 POLYVINYLCARBAZOLE
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open forms to the intensely colored closed isomers which could, in turn,

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(POLYVINYLCARBAZOLE OR POLYVINYLCARBAZOLES)
         15956 CARBAZOLE
         2128 CARBAZOLES
         16513 CARBAZOLE
                 (CARBAZOLE OR CARBAZOLES)
          5767 VINYLCARBAZOLE
           122 VINYLCARBAZOLES .
          5783 VINYLCARBAZOLE
                 (VINYLCARBAZOLE OR VINYLCARBAZOLES)
          1159 . PVK
            ·5 PVKS
          1161 PVK
                 (PVK OR PVKS)
           129 PVCZ
             8 L1 AND (((CHARGE OR HOLE)(2W)(TRANSPORT OR GENERAT?)) OR POLYVIN
               YLCARBAZOLE OR CARBAZOLE OR VINYLCARBAZOLE OR PVK OR PVCZ)
=> d all 1-18
    ANSWER 1 OF 8 CAPLUS COPYRIGHT 2005 ACS on STN
     2003:983002 CAPLUS
     140:154990
    Entered STN: 17 Dec 2003
    Electrical carrier-injection and transport characteristics of photochromic
       ***diarylethene***
                            films
     Tsujioka, Tsuyoshi; Masuda, Kazuya
    Department of Arts and Sciences, Osaka Kyoiku University, Kashiwara,
     Osaka, 582-8582, Japan
     Applied Physics Letters (2003), 83(24), 4978-4980
     CODEN: APPLAB; ISSN: 0003-6951
    American Institute of Physics
    Journal
    English
     76-1 (Electric Phenomena)
     Section cross-reference(s): 73
    Elec. carrier-injection and transport characteristics of photochromic
                           thin films were studied. Injected and transported
       ***diarylethene***
     carriers (injected current) to the ***diarylethene*** film were
     increased with the existence ratio of the closed-ring form, which was one
     isomerization state of the ***diarylethene*** mol. A dramatic
     increase was obsd. at a 12-percent existence ratio of the closed-ring
     mols. A lowering effect of the potential barrier, which is caused by a
     Poole-Frenkel-like effect with a short distance interaction potential for
    hole transportation by increasing the existence ratio of the closed-ring
    mols. is proposed as the mechanism behind this phenomenon.
       ***hole***
                     ***transport***
                                      photochromic arylethene film current
     injection
    Conduction electrons
    Electric current
     Electric current-potential relationship
     Electrooptical absorption
    Hole mobility
         ***Hole***
                       ***transport***
    Hole traps
     Isomers
     Poole-Frenkel effect
     Potential barrier
        (elec. carrier-injection and transport characteristics of photochromic
          ***diarylethene***
                               films)
    Isomerization
        (photoisomerization, UV; elec. carrier-injection and transport
        characteristics of photochromic
                                         ***diarylethene***
     406727-10-4
    RL: CPS (Chemical process); DEV (Device component use); PEP (Physical,
    engineering or chemical process); PRP (Properties); PYP (Physical
    process); PROC (Process); USES (Uses)
        (elec. carrier-injection and transport characteristics of photochromic
          ***diarylethene***
                               films)
    123847-85-8, NPB
    RL: DEV (Device component use); PEP (Physical, engineering or chemical
    process); PYP (Physical process); PROC (Process); USES (Uses)
        (.alpha.-NPB; elec. carrier-injection and transport characteristics of
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photochromic
                      ***diarylethene***
                                            films)
RE.CNT 9
             THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE
(1) Chidichimo, G; J Appl Phys 2001, V90, P4906 CAPLUS
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(4) Irie, M; Nature (London) 2002, V420, P759 CAPLUS
(5) Rodriguez, A; Appl Phys Lett 2001, V79, P461 CAPLUS
(6) Tsujioka, T; Appl Phys Lett 2001, V78, P2282 CAPLUS
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    ANSWER 2 OF 8 CAPLUS COPYRIGHT 2005 ACS on STN
L3
AN
    2003:594726 CAPLUS
DN
    139:283250
ED
    Entered STN: 04 Aug 2003
ΤI
    Organic bistable molecular memory using photochromic ***diarylethene***
ΑU
    Tsujioka, Tsuyoshi; Kondo, Hayato
    Faculty of Education, Department of Arts and Sciences, Osaka Kyoiku
CS
    University, Asahigaoka 4-698-1, Kashiwara, Osaka, 582-8582, Japan
    Applied Physics Letters (2003), 83(5), 937-939
SO
    CODEN: APPLAB; ISSN: 0003-6951
    American Institute of Physics
PB
DT
    Journal
LΑ
    English
    74-12 (Radiation Chemistry, Photochemistry, and Photographic and Other
CC
    Reprographic Processes)
    Section cross-reference(s): 76
    A principle of org. memory device using a bistable photochromic mol. is
AB
    presented that allows extremely high bit densities and very low power
     consumption. This device is based on an isomerization reaction of
    photochromic
                   ***diarylethene***
                                        mol. via its excited state by an elec.
     carrier injection, not by photon absorption. Exptl. data show that the
     reversible writing and nondestructive reading of information by the
     carrier injection is feasible. The advantages and properties of such an
    org. semiconductor memory using a bistable mol. are discussed.
st
    org bistable mol memory photochromic ***diarylethene***
     isomerization; elec excitation photochromic
                                                  ***diarylethene***
    bistable mol memory
IT
    Memory devices
        (nonvolatile; org. memory device using bistable photochromic
          ***diarylethene***
                              and based on isomerization of
                                                               ***diarylethene***
       via its excited state induced by elec. carrier injection)
IT
    Cyclization
    Photochromic materials
    Ring opening
        (org. memory device using bistable photochromic ***diarylethene***
       and based on isomerization of
                                      ***diarylethene*** via its excited
        state induced by elec. carrier injection)
IT
    50926-11-9, ITO
    RL: DEV (Device component use); USES (Uses)
        (anode; org. memory device using bistable photochromic
          ***diarylethene*** and based on isomerization of
                                                               ***diarylethene***
       via its excited state induced by elec. carrier injection)
     221042-24-6
    RL: DEV (Device component use); USES (Uses)
        (cathode; org. memory device using bistable photochromic
          ***diarylethene***
                              and based on isomerization of
                                                               ***diarylethene***
       via its excited state induced by elec. carrier injection)
    185690-39-5, 4,4',4''-Tris[N-(1-naphthyl)-N-phenylamino]-triphenylamine
    RL: DEV (Device component use); USES (Uses)
        (hole injection layer; org. memory device using bistable photochromic
          ***diarylethene*** and based on isomerization of ***diarylethene***
       via its excited state induced by elec. carrier injection)
    123847-85-8
    RL: DEV (Device component use); USES (Uses)
          ***hole***
                         ***transport***
                                          ; org. memory device using bistable
                      ***diarylethene***
                                          and based on isomerization of
         ***diarylethene*** via its excited state induced by elec. carrier
       injection)
    606127-31-5
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RL: CPS (Chemical process); DEV (Device component use); PEP (Physical,
    engineering or chemical process); PROC (Process); USES (Uses)
        (open form; org. memory device using bistable photochromic
          ***diarylethene*** and based on isomerization of ***diarylethene***
        via its excited state induced by elec. carrier injection)
     606127-30-4
     RL: CPS (Chemical process); DEV (Device component use); PEP (Physical,
     engineering or chemical process); PROC (Process); USES (Uses)
        (org. memory device using bistable photochromic ***diarylethene***
                                        ***diarylethene***
        and based on isomerization of
                                                             via its excited
        state induced by elec. carrier injection)
              THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
       13
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(7) Irie, M; Nature (London) 2002, V420, P759 CAPLUS
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(12) Taniguchi, A; Jpn J Appl Phys, Part 1 2001, V40, P7029 CAPLUS
(13) Tsujioka, T; Appl Phys Lett 2001, V78, P2282 CAPLUS
     ANSWER 3 OF 8 CAPLUS COPYRIGHT 2005 ACS on STN
     2002:6078 CAPLUS
     136:286972
     Entered STN: 03 Jan 2002
     Carrier injection/transport characteristics of photochromic
       ***diarylethene***
                            film
     Taniguchi, Akira; Tsujioka, Tsuyoshi; Hamada, Yuji; Shibata, Kenichi;
     Fuyuki, Takashi
     Graduate School of Materials Science, Nara Institute of Science and
     Technology, Ikoma, Nara, 630-0101, Japan
     Japanese Journal of Applied Physics, Part 1: Regular Papers, Short Notes &
     Review Papers (2001), 40(12), 7029-7030
     CODEN: JAPNDE
     Japan Society of Applied Physics
     Journal
     English
     76-1 (Electric Phenomena)
     Section cross-reference(s): 73
                    ***diarylethenes***
     Photochromic
                                          with elec. carrier (hole) controlling
     ability for nondestructive readout were investigated. The ionization
     potential of the photostationary colored ***diarylethene*** films was
     around 5.7-5.8 eV irresp. of differences in their mol. structures. Good
       ***hole***
                    injection/ ***transport*** characteristics were obsd. for
                            ***diarylethenes***
                                                  with triphenylamine groups.
     the colored films of
                         ***diarylethene***
     carrier transport
     Photochromic materials
        (carrier injection/transport characteristics of photochromic
          ***diarylethene***
                               film)
     Electric current carriers
     Hole (electron)
        (injection; carrier injection/transport characteristics of photochromic
          ***diarylethene***
                               film)
     Electric current carriers
        (transport; carrier injection/transport characteristics of photochromic
          ***diarylethene***
                              film)
                   314075-73-5
                                 406727-09-1
                                                             406727-11-5
     252253-50-2
                                               406727-10-4
     RL: PRP (Properties)
        (carrier injection/transport characteristics of photochromic
          ***diarylethene***
                              film)
RE.CNT
              THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD
(1) Adachi, C; Appl Phys Lett 1995, V66, P2679 CAPLUS
(2) Anon; Photo-Reactive Materials for Ultrahigh Density Optical Memory 1994
(3) Fukudome, M; Chem Lett 2001, V1, P70
(4) Hamano, M; Jpn J Appl Phys 1996, V35, P1764 CAPLUS
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(6) Irie, M; Chem Rev 2000, V100, P1685 CAPLUS
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    ANSWER 4 OF 8 CAPLUS COPYRIGHT 2005 ACS on STN
L3
AN
    2001:932566 CAPLUS
DN
    136:54934
ED
    Entered STN: 27 Dec 2001
ΤI
    Radiation-sensitive compositions containing photochromic compounds and
    thermally stable dosimeters manufactured from them
    Irie, Masahiro; Irie, Setsuko; Tanaka, Yuki
IN
PA
    Mitsubishi Chemical Corp., Japan
SO
    Jpn. Kokai Tokkyo Koho, 8 pp.
    CODEN: JKXXAF
DT
    Patent
LA
    Japanese
IC
    ICM C08L101-00
    ICS C08F012-02; C08F024-00; C08F026-06; C08F026-12; C08F028-06;
         C08F212-06; C08F212-08; C08K005-01; C08L025-00; C08L025-04;
         C08L025-16; C08L037-00; C08L039-04; C08L041-00; C09K009-02;
         G01T001-04
    38-3 (Plastics Fabrication and Uses)
    Section cross-reference(s): 25, 71
FAN.CNT 1
                                      APPLICATION NO. DATE
                       KIND
    PATENT NO.
                              DATE
                                         ______
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                              -----
                                                                -----
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                              20011225 JP 2000-178581 20000614
    JP 2001354862
                       A2
ΡI
                              20000614
PRAI JP 2000-178581
CLASS
            CLASS PATENT FAMILY CLASSIFICATION CODES
 PATENT NO.
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               ----
 JP 2001354862 ICM
                       C08L101-00
             . ICS
                       C08F012-02; C08F024-00; C08F026-06; C08F026-12;
                       C08F028-06; C08F212-06; C08F212-08; C08K005-01;
                       C08L025-00; C08L025-04; C08L025-16; C08L037-00;
                       C08L039-04; C08L041-00; C09K009-02; G01T001-04
AΒ
    The compns. contain thermally irreversible photochromic compds. and
    compds. having condensed polycyclic arom. structures. Thus, a transparent
     film manufd. from 0.90 g styrene-2-vinylnaphthalene copolymer and 0.10 g
     1,2-bis(2-methyl-5-phenyl-3-thienyl) ***perfluorocyclopentene***
    blue (600 nm absorption increase 0.070) after .gamma.-ray irradn. and no
    absorption change after 1 day at a room temp.
     radiation sensitivity thermally irreversible photochromic compd; dosimeter
ST
       ***diarylethene*** polycyclic arom compd naphthalene; vinylnaphthalene
     copolymer arylethene film thermal stability
     Polycyclic compounds
     RL: TEM (Technical or engineered material use); USES (Uses)
        (arom.; radiation-sensitive compns. contg. photochromic compns. and
       vinylnaphthalene copolymers or condensed polycyclic arom. compds. for
       dosimeters)
IT
    Aromatic compounds
     RL: TEM (Technical or engineered material use); USES (Uses)
        (polycyclic; radiation-sensitive compns. contg. photochromic compns.
       and vinylnaphthalene copolymers or condensed polycyclic arom. compds.
       for dosimeters)
IT
    Dosimeters
        (radiation-sensitive compns. contq. photochromic compns. and
       vinylnaphthalene copolymers or condensed polycyclic arom. compds. for
       dosimeters)
IT
     Photochromic materials
        (thermally irreversible; radiation-sensitive compns. contq.
       photochromic compns. and vinylnaphthalene copolymers or condensed
       polycyclic arom. compds. for dosimeters)
IT
     182003-69-6
     RL: PRP (Properties); TEM (Technical or engineered material use); USES
        (radiation-sensitive compns. contg. photochromic compns. and
       vinylnaphthalene copolymers or condensed polycyclic arom. compds. for
       dosimeters)
     85-01-8, Phenanthrene, uses
                                86-73-7, Fluorene
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95-15-8, Benzothiophene
                             119-65-3, Isoquinoline
                                                       120-12-7, Anthracene,
            120-72-9, Indole, uses
                                     129-00-0, Pyrene, uses
                                                              198-55-0,
     Perylene
                208-96-8, Acenaphthylene
                                          217-59-4, Triphenylene
                                                                    218-01-9,
                                           271-89-6, Benzofuran
     Chrysene
                229-87-8, Phenanthridine
                                                                  275-51-4,
     Azulene
               60279-76-7, Styrene-2-vinylnaphthalene copolymer
     RL: TEM (Technical or engineered material use); USES (Uses)
        (radiation-sensitive compns. contg. photochromic compns. and
        vinylnaphthalene copolymers or condensed polycyclic arom. compds. for
        dosimeters)
    ANSWER 5 OF 8 CAPLUS COPYRIGHT 2005 ACS on STN
     2000:764173 CAPLUS
     134:63722
     Entered STN: 01 Nov 2000
    Radiation sensitivity of photochromic
                                             ***diarylethenes***
     Irie, S.; Irie, M.
     Research Institute for Advanced Science and Technology, Osaka Prefecture
     University, Osaka, 599-8570, Japan
    Molecular Crystals and Liquid Crystals Science and Technology, Section A:
    Molecular Crystals and Liquid Crystals (2000), 345, 179-184
     CODEN: MCLCE9; ISSN: 1058-725X
     Gordon & Breach Science Publishers
     Journal
     English
     74-1 (Radiation Chemistry, Photochemistry, and Photographic and Other
     Reprographic Processes)
     Section cross-reference(s): 71
     The radiation sensitivity of photochromic dithienylethenes was studied
     with the aim of developing a new reusable color film dosimeter. Upon
     .gamma.-irradn., polystyrene films contg. 1,2-bis(2,5-dimethyl-3-thienyl)
       ***perfluorocyclopentene***
                                     (1) and 1,2-bis(2-methyl-5-phenyl-3-thienyl)
       ***perfluorocyclopentene***
                                     (2) turned red and blue, resp. The red and
     blue colors disappeared by visible-light irradn. and the films could be
     reused. In both films the absorption intensities of the colors linearly
     increased with the absorbed dose. From the color change it was possible
     to est. the radiation dose.
                                          ***diarylethene***
     radiation sensitivity photochromic
                                                               color dosimeter;
     gamma radiation color dosimeter photochromic
                                                    ***diarylethene***
     Dosimeters
        (gamma-ray, film; radiation sensitivity of photochromic
        dithienylethenes for development of reusable color dosimeter film)
     Polycarbonates, properties
     Polyolefins
     RL: PEP (Physical, engineering or chemical process); PRP (Properties);
     PROC (Process)
        (matrix; radiation sensitivity of photochromic dithienylethenes for
        development of reusable color dosimeter film)
     9003-53-6, Polystyrene
                              9011-14-7, PMMA
                                                25067-59-8, Poly(N-
       ***vinylcarbazole***
     RL: PEP (Physical, engineering or chemical process); PRP (Properties);
     PROC (Process)
        (matrix; radiation sensitivity of photochromic dithienylethenes for
        development of reusable color dosimeter film)
                  190394-25-3, 1,2-Bis(2,5-dimethyl-3-thienyl)
       ***perfluorocyclopentene***
                                       222972-69-2
                                                     277303-68-1
     RL: PEP (Physical, engineering or chemical process); PRP (Properties);
     PROC (Process)
        (radiation sensitivity of photochromic dithienylethenes for development
        of reusable color dosimeter film)
RE.CNT
              THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD
(1) Fujimura, R; Housyasen ni yoru Kotaigensyo to Senryosokutei 1985
(2) Irie, M; Bull Chem Soc Jpn 1998, V71, P985 CAPLUS
(3) Irie, M; Chem Lett 1995, P899 CAPLUS
(4) Irie, S; Bull Chem Soc Jpn 1999, V72, P1139 CAPLUS
(5) Kawanishi, S; Oyobutsuri 1982, V51, P293
(6) Yamagami, M; J Photopol Sci Tech 1995, V8, P101
    ANSWER 6 OF 8 CAPLUS COPYRIGHT 2005 ACS on STN
     2000:751164 CAPLUS
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91-20-3, Naphthalene, uses

91-22-5, Quinoline, uses

\*\*\*Carbazole\*\*\*

L3

AN

DN

ED

ТT ΑU

CS

SO

PB

DT

LA

CC

AΒ

IT

IT

L3

AN

DN

134:63711

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Entered STN: 25 Oct 2000
ED
    .Radiation-Induced Coloration of Photochromic Dithienylethene Derivatives
TI
     in Polymer Matrixes
AII
     Irie, Setsuko; Irie, Masahiro
     Res. Inst. Adv. Sci. Technol., Osaka Prefecture University, Sakai, Osaka,
CS
     599-8570, Japan
     Bulletin of the Chemical Society of Japan (2000), 73(10), 2385-2388
SO
     CODEN: BCSJA8; ISSN: 0009-2673
PB
     Chemical Society of Japan
DT
     Journal
     English
LA
     74-1 (Radiation Chemistry, Photochemistry, and Photographic and Other
CC
     Reprographic Processes)
     Section cross-reference(s): 71
AB
     The high-energy radiation-induced coloration of photochromic
     dithienylethenes was studied in various polymer matrixes with the aim of
     developing a reusable color plastic dosimeter. Upon .gamma.-irradn.,
     polystyrene films contg. 1,2-bis(2,5-dimethyl-3-thienyl)
       ***perfluorocyclopentene***
                                     1a and 1,2-bis(2-methyl-5-phenyl-3-thienyl)
       ***perfluorocyclopentene***
                                     2a turned red and blue, resp. The red and
     blue colors disappeared upon irradn. with visible light, and the films
     could be reused. In both films the absorption intensities of the colors
     increased linearly with the absorbed dose. Although radiation-induced
     coloration was clearly detected in polystyrene films, a color change was
     scarcely obsd. in poly(Me methacrylate) (PMMA) and poly(N-vinyl
       ***carbazole*** ) contg. 1a and 2a. Excitation energy transfer from
     polymers to dithienylethenes is considered to play a role in the
     coloration process in polymer films.
     gamma ray coloration photochromic dithienylethene polymer matrix film
ST
     dosimeter
IT
     Dosimeters
        (gamma-ray; radiation-induced coloration of photochromic
        dithienylethenes in relation to design of reusable color plastic
        dosimeters)
IT
     Polycarbonates, properties
     RL: PEP (Physical, engineering or chemical process); PRP (Properties);
        (matrix; radiation-induced coloration of photochromic dithienylethenes
        in relation to design of reusable color plastic dosimeters)
IT
     Energy transfer
     Optical absorption
     Photochromic materials
     Singlet state excitation
        (radiation-induced coloration of photochromic dithienylethenes in
        relation to design of reusable color plastic dosimeters)
     9003-53-6, Polystyrene
                             9011-14-7, PMMA
                                               25067-59-8, Poly(N-
IT
       ***vinylcarbazole***
     RL: PEP (Physical, engineering or chemical process); PRP (Properties);
     PROC (Process)
        (matrix; radiation-induced coloration of photochromic dithienylethenes
        in relation to design of reusable color plastic dosimeters)
                   313947-80-7
IT
     RL: FMU (Formation, unclassified); PEP (Physical, engineering or chemical
     process); PRP (Properties); FORM (Formation, nonpreparative); PROC
     (Process)
        (radiation-induced coloration of photochromic dithienylethenes in
        relation to design of reusable color plastic dosimeters)
                   190394-25-3, 1,2-Bis(2,5-dimethyl-3-thienyl)
       ***perfluorocyclopentene***
     RL: PEP (Physical, engineering or chemical process); PRP (Properties);
     PROC (Process)
        (radiation-induced coloration of photochromic dithienylethenes in
        relation to design of reusable color plastic dosimeters)
              THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
(1) Birks, J; Photophysics of Aromatic Molecules 1970
(2) Fujimura, R; Housyasen niyoru Kotaigensyou to Senryousokutei 1985
(3) Irie, M; Bull Chem Soc Jpn 1998, V71, P985 CAPLUS
(4) Irie, M; Chem Lett 1995, P899 CAPLUS
(5) Irie, M; J Org Chem 1995, V60, P8305 CAPLUS
(6) Irie, S; Bull Chem Soc Jpn 1999, V72, P1139 CAPLUS
(7) Johnson, G; J Chem Phys 1975, V62, P4697 CAPLUS
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(8) Kawanishi, S; Oyobuturi 1982, V51, P293
(9) Flopffer, W; J Polym Sci, Symp 1973, V40, P43
(10) Miyasaka, H; Chem Phys Lett 1997, V269, P281 CAPLUS
(11) Whitaker, B; Manual on Radiation Dosimetry 1970, P363
(12) Yamagami, M; J Photopol Sci Tech 1995, V8, P101
    ANSWER 7 OF 8 CAPLUS COPYRIGHT 2005 ACS on STN
L3
AN
    1998:785488 CAPLUS
DN
    130:160578
ED
    Entered STN: 15 Dec 1998
    Characterization of dye-doped PMMA/ ***PVK***
                                                      films as recording
TΙ
    materials
    Lafond, Christophe; Lessard, Roger A.; Bolte, Michele; Petkov, Ivan
ΑU
    Centre d'Optique Photonique et Laser (COPL), Universite Laval, Quebec,
CS
    G1K7P4, Can.
    Proceedings of SPIE-The International Society for Optical Engineering
SO
     (1998), 3417 (Photopolymer Device Physics, Chemistry and Applications IV),
     216-227
    CODEN: PSISDG; ISSN: 0277-786X
PB
    SPIE-The International Society for Optical Engineering
DT
    Journal
LA
    English
    74-12 (Radiation Chemistry, Photochemistry, and Photographic and Other
CC
    Reprographic Processes)
    The photoinduced reversible color change in photochromic mol. doped PMMA
AΒ
     and poly( ***vinylcarbazole*** ) ( ***PVK*** ) films was investigated.
    Upon UV and visible irradn., changes of an optical absorption of a closed
     form of a photochromic mol., in the beginning of the photoreaction,
     followed first-order kinetic and, then deviated from this order. The
     rate-consts. kUV of the coloring process, and kVIS of the bleaching
                                        ***diarylethenes***
                                                             studied, kUV, was
    process, were detd. For the two
     much larger in both polymer matrixes. However, kVIS was stronger in
       ***DVK***
                   for fulgide Aberchrome 670. Both rates were identical in
           Photochem. fatigue resistance was studied. For fulgide Aberchrome
     670 in PMMA matrix, the authors found 13% disappearance after 10 repeated
     UV and visible cycles.
    optical recording photochromic compd doped polymer film; photolysis
    photochromism fulgide doped polymer; ***diarylethene***
                                                                photochromism
     fulgide doped polymer
     UV and visible spectra
        (absorption; photoinduced reversible color changes in PMMA and poly(
          ***vinylcarbazole*** ) films contg. fulgide or   ***diarylethene***
       photochromic mols.)
     Absorption spectra
     Optical recording
     Photochromic materials
     Photochromism
     Photolysis
     Photolysis kinetics
        (photoinduced reversible color changes in PMMA and poly(
          ***vinylcarbazole***  ) films contg. fulgide or   ***diarylethene***
       photochromic mols.)
IT
     Isomerization
        (photoisomerization; photoinduced reversible color changes in PMMA and
        poly( ***vinylcarbazole*** ) films contg. fulgide or
          ***diarylethene***
                             photochromic mols.)
                      25067-59-8, Poly( ***vinylcarbazole*** )
     9011-14-7, PMMA
                   112440-48-9
                                 142381-71-3
                                               220191-33-3
                                                            220191-36-6
     112440-47-8
     220191-38-8
     RL: PEP (Physical, engineering or chemical process); PRP (Properties);
     PROC (Process)
        (photoinduced reversible color changes in PMMA and poly(
          ***vinylcarbazole*** ) films contg. fulgide or
                                                           ***diarylethene***
        photochromic mols.)
              THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
       10
(1) Durr, H; Photochromism: Molecules and Systems 1990, P5
(2) Ghailane, F; Ph D thesis Universite Laval 1995
(3) Hanazawa, M; J Chem Soc Chem Commun 1992, V3, P206
(4) Hoshino, M; J Photochem Photobiol A: Chem 1997, V105(1), P75 CAPLUS
(5) Irie, M; J Org Chem 1988, V53, P803 CAPLUS
(6) Kaneko, A; Bull Chem Soc Jpn 1988, V61(10), P3569 CAPLUS
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(7) Tsuyioka, T; J Photochem Photobiol A: Chem 1997, V104, P203
(8) Whitall, J; Photochromism: Molecules and Systems 1990, P467
(9) Yokoyama, Y; Bull Chem Soc Jpn 1990, V63(6), P1607 CAPLUS
(10) Yokoyama, Y; J Chem Soc Chem Commun 1991, V24, P1722
     ANSWER 8 OF 8 CAPLUS COPYRIGHT 2005 ACS on STN
L3
     1997:457764 CAPLUS
AN
     127:115179
DN
ED
     Entered STN: 23 Jul 1997
     Charge injection photocontrol at the metal/organic interface by a
     photochromic compound
     Honma, Hisakazu; Yokoyama, Masaaki
ΑU
    Material and Life Science, Grad. School Engineering, Osaka Univ., Suita,
CS
     565, Japan
     Denshi Shashin Gakkaishi (1997), 36(1), 5-10
SO
     CODEN: DSHGDD; ISSN: 0387-916X
    Denshi Shashin Gakkai
PB
DT
    Journal
LA
     Japanese
     74-12 (Radiation Chemistry, Photochemistry, and Photographic and Other
CC
    Reprographic Processes)
AΒ
       ***Diarylethenes***
                            are known as thermally stable photochromic
     compds., particularly of which
                                     ***diarylethene***
                                                          derivs. having
    bithiophene as the side groups undergo the great change in the
     .pi.-electron conjugation length between the open-form and closed-form
    upon their photo-isomerization. Consequently, the ionization potential
     (Ip) is expected to change between both forms. In this paper, we have
     applied the difference of Ip between these two forms for controlling the
    hole injection from a metal electrode to an org. film. We have succeeded
     in demonstrating the photo-switch of the injection current on the order of
     sub-miliampare upon photo-isomerization by inserting a thin film of
       ***diarylethene*** deriv. in-between metal and org.
                                                               ***charge***
       ***transport*** layer.
    photochromic
                   ***diarylethene*** charge injection photocontrol
ST
    photoswitch
    Electrophotographic photoconductors (photoreceptors)
IT
    Optical recording materials
    Optical switching
     Photochromic materials
     Spatial light modulators
        (charge injection photocontrol at the metal/org. interface by a
       photochromic compd.)
TΤ
     Isomerization
        (photoisomerization; charge injection photocontrol at the metal/org.
        interface by a photochromic compd.)
     159590-09-7P
IT
    RL: DEV (Device component use); PNU (Preparation, unclassified); PRP
     (Properties); PREP (Preparation); USES (Uses)
        (charge injection photocontrol at the metal/org. interface by a
       photochromic compd.)
=> d his
     (FILE 'HOME' ENTERED AT 10:05:32 ON 14 NOV 2005)
    FILE 'CAPLUS' ENTERED AT 10:05:38 ON 14 NOV 2005
L1
            828 S (DIARYLETHENE OR PERFLUOROCYCLOPENTENE)
L2
             14 S L1 AND (NLO OR NONLINEAR OR HYPERPOLARIZ? OR NON(2A)LINEAR)
              8 S L1 AND (((CHARGE OR HOLE)(2W)(TRANSPORT OR GENERAT?)) OR POLY
L3
=> log y
COST IN U.S. DOLLARS
                                                 SINCE FILE
                                                                  TOTAL
                                                      ENTRY
                                                               SESSION
FULL ESTIMATED COST
                                                      97.83
                                                                  98.04
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)
                                                 SINCE FILE
                                                                 TOTAL
                                                      ENTRY
                                                               SESSION
CA SUBSCRIBER PRICE
                                                     -16.06
                                                                -16.06
```

Ref #	Hits	Search Query	DBs	Default Operator	Plurals	Time Stamp
L1	45473 ·	(photochromic or spiropyran\$1 or fulgide\$4 or stilbene\$1 or leuco or diarylethane\$\$ or diheteroarylethane\$2 or dithiophene\$7)	US-PGPUB; USPAT; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2005/11/14 08:21
L2	171897	(photorefract\$6 or polyvinylcarbazole\$1 or carbazole\$1 or vinylcarbazole\$1 or ((charge or hole) near5 (transport or generat\$6)))	US-PGPUB; USPAT; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2005/11/14 08:21
L3	2272	l1 same l2	US-PGPUB; USPAT; EPO; JPO; DERWENT; IBM_TDB	OR	ON '	2005/11/14 08:21
L4	171934	(photorefract\$6 or pvcz or polyvinylcarbazole\$1 or carbazole\$1 or vinylcarbazole\$1 or ((charge or hole) near5 (transport or generat\$6)))	US-PGPUB; USPAT; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2005/11/14 08:32
L5	2272	l1 same l4	US-PGPUB; USPAT; EPO; JPO; DERWENT; IBM_TDB	OR	ON .	2005/11/14 08:21
L6	7	(diarylethane\$\$ or diheteroarylethane\$2 or dithiophene\$7) same I4	US-PGPUB; USPAT; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2005/11/14 08:23
L7	7	(diarylethane\$\$ or diheteroarylethane\$2 or dithiophene\$7) and I3	US-PGPUB; USPAT; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2005/11/14 08:23
L8		(diarylethane\$\$ or diheteroarylethane\$2 or dithiophene\$7) and I4	US-PGPUB; USPAT; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2005/11/14 08:24
L9	<b>75</b>	l8 and @ad<"20030818"	US-PGPUB; USPAT; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2005/11/14 08:33
L10	325283	(ito or ((indium or tin) near5 oxide\$1))	US-PGPUB; USPAT; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2005/11/14 08:31

L11	517387	l10 or (l1 or l4)	US-PGPUB; USPAT; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2005/11/14 08:31
L12	3706	I10 same (I1 or I4)	US-PGPUB; USPAT; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2005/11/14 08:31
L13	410	I5 and I12	US-PGPUB; USPAT; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2005/11/14 08:32
L14	410	(pvcz or polyvinylcarbazole\$1 or carbazole\$1 or vinylcarbazole\$1 or ((charge or hole) near5 (transport or generat\$6))) and l13	US-PGPUB; USPAT; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2005/11/14 08:32
L15	328	I13 and @ad<"20030818"	US-PGPUB; USPAT; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2005/11/14 08:33
L16	136	I15 not (tonor or Toner or electrophoto\$8 or emitting).ti,ab.	US-PGPUB; USPAT; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2005/11/14 08:37
L17	288864	(nonlinear or hyperpolariz\$6 or (second near5 order) or nlo)	US-PGPUB; USPAT; EPO; JPO; DERWENT; IBM_TDB	OR	ON ·	2005/11/14 08:38
L18	27	l13 and l17	US-PGPUB; USPAT; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2005/11/14 08:38